BRATTLEBORO MEMORIAL HOSPITAL

2018 LAB GUIDE GENERAL INFORMATION
ADMINISTRATION

Medical Director of Laboratory.......................... Christopher Appleton, D.O. FACP

Administrative Director of Laboratory Services........ Imogene Drakes, PhD, FACHE

Laboratory Supervisor......................................................... Deborah Gay, MT, ASCP
TELEPHONE / FAX NUMBERS

General Information....................................................................................257-8311

Results Inquiry:
                                    Clinical Fax.....................................................................................257-8287

Outpatient Phlebotomy..............................................................................275.3633
LABORATORY HOURS

LABORATORY – CUSTOMER SERVICE  MONDAY - FRIDAY  8AM – 4:30PM

PATHOLOGY AIDE  MONDAY – FRIDAY  9 AM – 3:30PM*

* All other hours – Contact Pathologist on-call through BMH Operator (802) 257-0341

OUTPATIENT HOURS

MONDAY – FRIDAY  7AM – 6PM
SATURDAY  8AM – 12 PM
SUNDAY  CLOSED
HOLIDAY HOURS  AS POSTED
HOW TO COMPLETE OUTPATIENT LABORATORY REQUISITIONS

All outpatient Laboratory tests must be ordered on a Laboratory requisition (see Main Laboratory requisition on following page).

Mandatory Information Required:

1. Patient’s full name, date of birth and gender
2. Patient’s address and phone number
3. Patient’s primary and secondary insurance information
4. Patient under 18 years, parent/guardian’s name and address and social security number
5. Authorization and assignment signature
6. Fully legible name of authorized HCP ordering test
7. Diagnosis / Symptoms / Medical necessity / ICD-10 code (Choose from selection on back page of requisition or write ICD-10 code(s) on lines at the top of the back page).
8. Test(s) ordered
9. Specimen description

**IT IS ALSO VERY IMPORTANT TO INCLUDE THE DATE AND TIME OF SPECIMEN COLLECTION** if you are collecting the specimen.

Additional Information

1. If you need a test done STAT, place an X in the STAT box on the front of the requisition.
2. If you need results called or faxed to your office, please record this on the requisition and supply the phone/fax number.
3. If another physician requires a copy of the laboratory report, please print the first and last name of the physician on the requisition in the “copy to” box.
4. If the information described above is not provided, a request will be made for a corrected requisition. Testing will be delayed until the appropriate information is provided.

NOTES:

For certain infectious diseases (e.g., malaria), travel or other risk factors should be listed. Requisitions can be obtained from the laboratory by calling (802) 257-8311.
CRITICAL LAB VALUES TO BE CALLED (3 Pages)

**CHEMISTRY SECTION**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bilirubin, Total (All ages)</td>
<td>≥ 13 mg/dL</td>
</tr>
<tr>
<td>Blood Urea Nitrogen</td>
<td>&gt; 104 mg/dL</td>
</tr>
<tr>
<td>CO2</td>
<td>&lt;10 mmol/L</td>
</tr>
<tr>
<td></td>
<td>&gt;40 mmol/L</td>
</tr>
<tr>
<td>Ca</td>
<td>&lt;7.0 mg/dL</td>
</tr>
<tr>
<td></td>
<td>&gt;14.0 mg/dL</td>
</tr>
<tr>
<td>Creatinine</td>
<td>&gt; 7.4 mg/dL</td>
</tr>
<tr>
<td>Glucose (&gt; 1 Month)</td>
<td>&lt;50 mg/dL</td>
</tr>
<tr>
<td></td>
<td>&gt;500 mg/dL</td>
</tr>
<tr>
<td>Glucose (Neonates)</td>
<td>&lt;40 mg/dL</td>
</tr>
<tr>
<td></td>
<td>&gt;200 mg/dL</td>
</tr>
<tr>
<td>K</td>
<td>&lt;3.0 mmol/L</td>
</tr>
<tr>
<td></td>
<td>&gt;6.0 mmol/L</td>
</tr>
<tr>
<td>Na</td>
<td>&lt;125 mmol/L</td>
</tr>
<tr>
<td></td>
<td>&gt;160 mmol/L</td>
</tr>
<tr>
<td>Mg</td>
<td>&lt;1.0 mg/dL</td>
</tr>
<tr>
<td></td>
<td>&gt;4.8 mg/dL</td>
</tr>
<tr>
<td>Phosphorus</td>
<td>&lt; 1.1 mg/dL</td>
</tr>
<tr>
<td>Troponin I</td>
<td>≥ 0.1 ng/mL</td>
</tr>
<tr>
<td>Vitamin D</td>
<td>≥100 ng/ml</td>
</tr>
</tbody>
</table>

**DRUG LEVELS GREATER THAN**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetaminophen</td>
<td>&gt;150 ug/mL</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>&gt; 30 ug/mL</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>&gt; 15 ug/mL</td>
</tr>
<tr>
<td>Salicylate</td>
<td>&gt;30 mg/dL</td>
</tr>
<tr>
<td>Digoxin</td>
<td>&gt;2.5 ng/mL</td>
</tr>
<tr>
<td>Theophylline</td>
<td>&gt;25 ug/mL</td>
</tr>
<tr>
<td>Gentamicin(peak)</td>
<td>&gt;12 ug/mL</td>
</tr>
<tr>
<td>Tobramycin(peak)</td>
<td>&gt;12 ug/mL</td>
</tr>
<tr>
<td>Gentamicin(random)</td>
<td>&gt;13 ug/mL</td>
</tr>
<tr>
<td>Tobramycin(trough)</td>
<td>&gt;2.0 ug/mL</td>
</tr>
<tr>
<td>Gentamicin(trough)</td>
<td>&gt;2.0 ug/mL</td>
</tr>
<tr>
<td>Valproic Acid</td>
<td>&gt;200 ug/mL</td>
</tr>
<tr>
<td>Lithium</td>
<td>&gt;1.6 mmol/L</td>
</tr>
<tr>
<td>Vancomycin (peak)</td>
<td>&gt;80 ug/mL</td>
</tr>
<tr>
<td>Valcomycin (trough)</td>
<td>&gt;25 ug/mL</td>
</tr>
</tbody>
</table>
HEMATOLOGY SECTION

<table>
<thead>
<tr>
<th>TEST</th>
<th>“low” critical value</th>
<th>“high” critical value</th>
</tr>
</thead>
<tbody>
<tr>
<td>WBC (Newborn)</td>
<td>&lt; 4.0 K/µL</td>
<td>&gt; 30.0 K/µL</td>
</tr>
<tr>
<td>WBC (Adult)</td>
<td>&lt; 1.0 K/µL</td>
<td>&gt; 20.0 K/µL</td>
</tr>
<tr>
<td>Platelets (Adult)</td>
<td>&lt; 40 K/µL</td>
<td>None</td>
</tr>
<tr>
<td>Blasts, Differential</td>
<td>Present</td>
<td>Present</td>
</tr>
<tr>
<td>Hemoglobin, (Newborn)</td>
<td>&lt; 9.7 ngm/L</td>
<td>&gt; 22.3 ngm/L</td>
</tr>
<tr>
<td>Hematocrit, (Newborn)</td>
<td>&lt; 29 %</td>
<td>&gt; 67 %</td>
</tr>
<tr>
<td>Hemoglobin, (Adult)</td>
<td>&lt; 8.0 ngm/L</td>
<td>&gt; 20.0 ngm/L</td>
</tr>
<tr>
<td>Hematocrit, (Adult)</td>
<td>&lt; 24%</td>
<td>&gt; 60 %</td>
</tr>
<tr>
<td>Heparin, Low Mol. Wt.</td>
<td>None</td>
<td>&gt; 2.0 IU/mL</td>
</tr>
<tr>
<td>Heparin, Unfractionated</td>
<td>None</td>
<td>&gt; 1.0 IU/mL</td>
</tr>
<tr>
<td>APTT</td>
<td>None</td>
<td>&gt; 119.0 seconds</td>
</tr>
<tr>
<td>Prot, (PT)- (Adult)</td>
<td>None</td>
<td>&gt; 44.3 seconds</td>
</tr>
<tr>
<td>PT INR</td>
<td></td>
<td>4.7</td>
</tr>
<tr>
<td>Neutrophil ABS % (auto Diff)</td>
<td>&lt; 0.49 K/µL</td>
<td></td>
</tr>
</tbody>
</table>

MICROBIOLOGY SECTION

<table>
<thead>
<tr>
<th>TEST</th>
<th>critical value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stool: Salmonella/Shigella</td>
<td>positive</td>
</tr>
<tr>
<td>E.Coli O157, E. coli (shigatoxin), Vibrio, Yersinia called by VT State</td>
<td>positive</td>
</tr>
<tr>
<td>CSF Smear and/or culture</td>
<td>positive</td>
</tr>
<tr>
<td>Rapid Strep Screen</td>
<td>positive</td>
</tr>
<tr>
<td>Acid Fast Smear and/or AFB Culture</td>
<td>positive</td>
</tr>
<tr>
<td>Blood Culture results</td>
<td>positive</td>
</tr>
<tr>
<td>MRSA on inpatients or Nursing Home</td>
<td>positive</td>
</tr>
<tr>
<td>VRE on inpatients or Nursing Home</td>
<td>positive</td>
</tr>
<tr>
<td>Group B Streptococci isolated from neonates or infants to age 3 months</td>
<td>All are called</td>
</tr>
<tr>
<td>B. Pertussis are called by VT State lab</td>
<td>positive</td>
</tr>
<tr>
<td>Fetal Fibronectin</td>
<td>positive</td>
</tr>
<tr>
<td>Giardia Antigen</td>
<td>positive</td>
</tr>
<tr>
<td>C. Difficile Toxin</td>
<td>positive</td>
</tr>
<tr>
<td>Chlamydia Trachomatis: Neisseria Gonorrhoeae Genprobes are called by UVMCC (Reference Lab)</td>
<td>positive</td>
</tr>
<tr>
<td>Legionella antigen screen</td>
<td>positive</td>
</tr>
<tr>
<td>Influenza A or B</td>
<td>positive</td>
</tr>
<tr>
<td>Campylobacter</td>
<td>positive</td>
</tr>
</tbody>
</table>
SEROLOGY SECTION

**CRITICAL VALUE – TEST**

<table>
<thead>
<tr>
<th>Test</th>
<th>Critical Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive - Herpes Simplex virus culture from any urogenital site of a woman of childbearing age (10-50 years of age) called by UVMMC (Reference Lab).</td>
<td></td>
</tr>
<tr>
<td>Positive - Viral culture from any site of a neonate called by UVMMC or MML (Reference Labs)</td>
<td></td>
</tr>
<tr>
<td>Positive – RSV</td>
<td></td>
</tr>
</tbody>
</table>

URINALYSIS SECTION:

<table>
<thead>
<tr>
<th>Test</th>
<th>Critical Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose (newborn)</td>
<td>Positive</td>
</tr>
<tr>
<td>Ketones (newborn)</td>
<td>Positive</td>
</tr>
<tr>
<td>Red blood cell casts</td>
<td>Positive</td>
</tr>
<tr>
<td>Reducing substance (&lt;1 month)</td>
<td>Positive</td>
</tr>
</tbody>
</table>

BLOOD BANK/TRANSFUSION SERVICE:

<table>
<thead>
<tr>
<th>Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incompatible crossmatch in setting of urgent blood need.</td>
</tr>
<tr>
<td>Transfusion reaction investigations showing a hemolytic reaction.</td>
</tr>
<tr>
<td>Unavailability of products to fill and order.</td>
</tr>
</tbody>
</table>

SURGICAL PATHOLOGY SECTION:

<table>
<thead>
<tr>
<th>Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Significant unsuspected diagnoses</td>
</tr>
<tr>
<td>Significant discrepancy between frozen and permanent diagnosis, with potential major impact on patient care</td>
</tr>
</tbody>
</table>

DEFINITIONS:

1) Licensed care giver: Refers to Physicians, Allied Health Staff, RN’s, or LPN’s.
2) Outpatient: This is any outpatient from a physician office, nursing home, or VNA at the time of the critical value report. Inpatient is any patient located on Med/Surg 2nd or 3rd floor, ACU, SCU, OR, Birth Center/ Nursery, Short Stay.

TRAINING:

All Technical and Phlebotomy staff are trained during orientation for new employees.
REFERENCE RANGE (NORMAL VALUES)

Reference ranges are guides rather than absolute indicators of health and disease. Values for healthy persons often overlap with values for persons afflicted with disease. Laboratory values may vary because of methodological differences and/or modes of standardization which exist between various laboratories.

Therapeutic and toxic drug ranges are those commonly accepted on the basis of current knowledge and recommended values of current reagent manufacturers.

REFERENCES:

4. Roche Package inserts for the respective reagents.
CRITICAL TESTS

Results will always be called to Provider. Expected turnaround time from time of specimen receipt in Lab to time of phoned results in parentheses.

- Frozen Sections (20 minutes)
- Troponin I (30 minutes)
CALLING CRITICAL RESULTS

PROCEDURE:

Once “critical” values for a test are established, laboratory personnel are required to follow this protocol for notification when critical results are obtained.

Call immediately to notify a licensed caregiver on the floor, in a physician’s Office, Nursing Home, or VNA patient.

a. Verify that patient is at this location (Nursing Home), or a patient at this office, or is being treated by VNA.

b. All Out Patient Critical Values” will be called after the lab test is verified and indicated in LIS that a “Critical Result Value has occurred. Critical values will be reported to a licensed Caregiver (s) caring for the patient within 60 minutes after the test is verified in LIS.

Lab staff will state to a caregiver that the results of a test are in a “Critical” value range and then give the critical value.

a) Lab will ask the care giver to repeat back to laboratory “the result or value reported.”

b) Lab staff will document the telephone call in the LIS in the “critical result area” place date, exact time, and the name of whom it was called to. This data will also appear in the various lab section reports.

Licensed caregiver in physician’s Office, Nursing Home, or VNA needs to immediately notify the “responsible licensed caregiver” who will act on the critical test result just being reported.

Critical test results may not be left on an answering machine. Please try several times to call. BMH Pathologist will be notified if the “Outpatient” ordering physician/licensed caregiver could not be reached.

Other results may be deemed critical if, in the opinion of the Pathologist or Technologist, the results may indicate the patient may require urgent care.

Note: Laboratory technical staff are responsible for notifying ordering personnel concerning turnaround time delays due to processing issues such as a dilution due to a “high” out of range message from a lab analyzer, instrument malfunction, quality control problems, extended processing, workload backup, etc., and assuring them that they will receive test results ASAP.
CRITERIA FOR ACCEPTABLE SPECIMENS

Specimens can be accepted and tested, if they meet the following guidelines:

1. **LABELING** – all specimens and aliquots must be received with a label that contains:
   - the patient’s full name
   - at least one other unique identifier (i.e., medical record number or date of birth)
   - date and time of collection (acceptable if on requisition only)
   - identity of the individual collecting BMH drawn samples and all Blood Bank specimens
   - all aliquots must bear the identity of the individual preparing the aliquot

   All Blood Bank samples for cross-match or type and screen must be labeled using the Secureline identification system.

2. **OUTPATIENT (OP) LABORATORY REQUISITION** – all OP specimens must be accompanied by a complete requisition. The form **must** contain the following:
   - Patient’s name
   - Patient’s sex
   - Patient’s date of birth
   - Name of physician or person legally authorized to order testing
   - Tests requested
   - Diagnosis (ICD 10) appropriate for all tests ordered.
   - Time and date of specimen collection, when specimen is accompanying requisition
   - Source of specimen, when appropriate
   - Clinical information, when appropriate
   - Completed consent form, when appropriate

3. **SPECIMEN CONTAINER** – the exterior must be intact and free of contamination by blood or body fluids. If the specimen is contained in a syringe, the needle must have been removed and replaced with a firmly sealed cap.

4. **VOLUME OF SPECIMEN** – the appropriate volume of specimen must be collected to meet testing requirements.

5. **COLLECTION DEVICE/PRESERVATIVE** – specimens must be submitted in the proper collection device and with the correct preservative.
CRITERIA FOR REJECTING SPECIMENS

Any specimen arriving in the laboratory that fails to meet criteria will be withheld from analysis until the deficiency has been resolved.

**Never** discard any rejected specimen before its normal discard date (See Add-on or Storage Requirements for each test in the Test Menu) or 72 hours. Whenever possible, a replacement specimen should be obtained. If one cannot be obtained, the clinician must be notified.

Unstable specimens/analytes or unique samples that cannot be recollected may need to be accepted even though the specimen is suboptimal. These specimens would include:
- CSF or other body fluids
- capillary/fingerstick specimens
- cord blood samples
- tissues
- culture specimens obtained prior to initiation of antibiotic therapy
- pediatric nasopharyngeal washings
- Pap smears

The original sample cannot be relabeled. All rejected specimens are to be retained in a designated area of the refrigerator for as long as their discard dates.

In the event that a specimen is unstable, unique or cannot be recollected, the physician and the individual who collected the sample (if not the same person) must certify in writing that (1) the specimen is irreplaceable and (2) the correct specimen information is accurate before the sample can be processed. (See example of the Authorization to Test Irretrievable Specimen on the next page). If the sample was mislabeled, the incorrect label cannot be removed. The correct LIS label can be placed on the sample. It is imperative that technical staff be made aware of the labeling discrepancy. Following the completion of the requested assay, technical staff must add a disclaimer to the results by appending a comment such as Specimen labeling issue, assay performed at the request of the Healthcare Provider).

Documentation in the LIS must occur whenever the Authorization to Test Irretrievable Specimen (see next page) is used to accept an unlabeled or mislabeled specimen. The signed form will be retained in the laboratory and an Incident Report must be completed.
AUTHORIZATION TO TEST IRRETRIEVABLE SPECIMEN

Name of Patient ____________________                           Date of Birth ________________

Medical Record Number ____________________  Accession Number _______________

Date Specimen Collected ________________.

Ordering Provider ________________

Test Ordered ____________________________________________________

Type of Specimen ________________________________________________

Source of Specimen ____________________________________________

Signature of Provider authorizing testing of specimen ______________Date ______

Laboratory Medical Director’s signature ____________________________Date _____

Form Created 10/29/2015
PATIENT IDENTIFICATION

The phlebotomist will use two patient identifiers before drawing blood:

**In an Outpatient setting:**

Ask the patient a direct question, “Can you give me your full name please?” and “What is your date of birth?” Compare the information stated by the patient with information on the computer labels or with the requisition slip.

Nursing home patients must also be identified using two unique identifiers. This is usually name and date of birth. If the patient/resident is unable to provide this information, it should be provided by a nursing home employee, unless a valid band is worn by the patient.

**In an Inpatient setting:**

Compare name and Medical Record # on the patient’s identification bracelet with that on your labels or requisition. This information must be **identical**! Usually the ID bracelet is on the patient’s wrist. In some cases, it may be on the patient’s ankle. Request a nurse to identify a patient who does not have an identification bracelet. A bracelet should be on the patient’s wrist except in cases when it is not feasible. In this case, have the nurse taking care of the patient identify the patient for you. Make a note on the requisition of the nurse who identified the patient.

Patient who is unconscious, too young, mentally incompetent, or does not speak the language of the phlebotomist:
Ask the nurse to identify the patient by name and Medical Record # or date of birth. Compare this data with the information on the request form. For outpatients, a relative or friend may be asked to identify the patient by name and date of birth.

Procedure for identifying Unidentified Emergency Patients:
The patient must be positively identified when the specimen is collected. The unidentified emergency patient is given a temporary designation until positive identification can be made. In all cases, the name and hospital number of the emergency identification are attached to the patient’s body either by wristband or some similar device.
REFERRAL OF SPECIMENS

Skin scrapings, conjunctival scrapings, throat swabs, Tzanck preparations, nasopharyngeal swabs, (e.g., for B. pertussis), and lavage for viruses are samples that are collected from procedures performed by the ordering clinic or providers on the floors.
SPECIMEN COLLECTION

The laboratory provides staff to assist in the collection of venous blood samples. Some nursing units collect their own samples and require less assistance. Other units rely solely on the laboratory. In either instance, the laboratory will respond to requests for assistance on either a scheduled or STAT basis as described below.

A. Blood collection schedules: There are five ways to request blood draws
   1. Early AM rounds are completed between 0600 - 0800.
   2. Periodic routine rounds may be scheduled after AM rounds, until 2300 and as needed during 3rd shift.
   3. Timed draws are scheduled as requested.
   4. STAT requests will be collected within 20 minutes of lab notification
   5. The majority of Laboratory tests are normally scheduled for early 6AM draws.

B. All specimens submitted to the Laboratory will be labeled in ink with:
   1. Patient’s full name
   2. Medical record number or date of birth
   3. Location
   4. Date collected
   5. Time collected
   6. Initials of collection personnel for all specimen types.
   7. Site / source of specimen as appropriate.

**NOTE:** All Inpatient specimens should be sent to the lab with the computer-generated label.

C. Specimens submitted on slides:
   1. Slide must be labeled in pencil.
   2. Slide must have patient’s full name and medical record number or date of birth.
   3. Slide container must be labeled with all the information listed in Item B above.

If the information described above is not provided, the specimen will be rejected and the nursing unit or other area initiating the request will be informed of the error. A request will be made for a corrected specimen. If specimens are not collected according to required procedures, a staff member will notify the nursing unit. If the patient is an outpatient, the attending physician’s office will be notified. Recommended collection procedure may be found listed in the individual test section. Hemolysis and/or lipemia-free specimens are required for certain procedures. If testing is performed on hemolyzed or lipemic specimens, a notation will appear on the report form. Any other observed interfering substance will also be noted on the report form.

☐ Specimen labeling must be on the actual sample container, not on an over wrap container or bag.
☐ Specimens must be submitted in solid sided, screw capped containers.
☐ Baggies are not acceptable.
SPECIMEN CONTAMINATION

Requisitions or other paper accidentally contaminated with specimens should be discarded into an appropriate container and a new requisition form made out. Specimen containers, whose external surface becomes contaminated, should be decontaminated in its entirety with an EPA-approved hospital disinfectant.

**Standard Precautions**
All specimens are presumed to be potentially infectious and are handled following “Standard Precautions.”
SUPPLIES FOR PHYSICIAN OFFICES

The BMH Laboratory provides a variety of collection supplies to assist clinicians in obtaining samples for testing in the BMH Lab and in the Reference Labs used by BMH. An order form for supplies may be obtained by calling the laboratory at 257-8311.

Urgent Inpatient supply orders will be filled within 24 hours and non-urgent Inpatient and Outpatient supply orders will be filled within 72 hours.

It is the understanding that supplies requested are used for the sole purpose of sending samples to the BMH Laboratory. As part of the federally mandated compliance program, the laboratory may periodically audit the relationship of supplies requested to specimens received.
BLOOD BANK SECTION (BB)

The Blood Bank is located in the Main Laboratory on the ground floor. The telephone is extension 8311. Transfusion services are available 24 hours per day, 7 days per week.

**Indications for Transfusion**

Blood product transfusions should only be given when medically necessary. Indications for transfusions/recommendations are embedded within printed transfusion orders and on electronic orders.
BB - EMERGENCY (UNCROSSMATCHED) BLOOD PROTOCOL

For an order of UNCROSSMATCHED blood, call the Blood Bank (Ext. 8311) indicating the need for Uncrossmatched blood, stating the number of units needed. This will give the Blood Bank Tech an opportunity to begin processing the necessary paperwork and deliver the blood to the appropriate unit.

The UNCROSSMATCHED blood will be issued with a fluorescent orange “UNCROSSMATCHED BLOOD” sticker attached to the face of the unit. The “Emergency Transfusion Request” form indicating the status of the bloods issued and listing the donor units will accompany the Uncrossmatched units of blood. The Physician MUST SIGN the request form and return the form to the Blood Bank Tech.

In cases of extreme emergency where there is not sufficient time to perform a blood type, O Negative packed cells will be released.
BB - FRESH FROZEN PLASMA

A call to the Blood Bank for Fresh Frozen Plasma (FFP) should be made 1 hour before the expected infusion (this is the time required to thaw the frozen plasma).

Fresh Frozen Plasma should be issued as ABO compatible (the Rh factor is insignificant). The FFP does not need to be crossmatched.

The sooner FFP is infused after being thawed, the greater the survival of the labile coagulation factors. FFP must be used within 24 hours after being thawed or it must be discarded (wasted).
BB - GUIDELINES FOR BLOOD PRODUCT TRANSFUSION

<table>
<thead>
<tr>
<th>PHYSICIAN'S ORDERS</th>
<th>BLOOD &amp; BLOOD PRODUCTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>NOTED (total)</td>
<td></td>
</tr>
<tr>
<td>G a 1 g 1 c 1</td>
<td></td>
</tr>
<tr>
<td>STAT</td>
<td></td>
</tr>
<tr>
<td>ROUTINE</td>
<td></td>
</tr>
<tr>
<td>Pre-op surg Date</td>
<td></td>
</tr>
<tr>
<td>Time</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>BLOOD PRODUCTS</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>INDICATION - CHECK ONE</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hemoglobin/Hematocrit</td>
</tr>
<tr>
<td></td>
<td>When to Transfuse - Guideline</td>
</tr>
<tr>
<td></td>
<td>□ &lt; 6 g/dL 18%</td>
</tr>
<tr>
<td></td>
<td>Almost always</td>
</tr>
<tr>
<td></td>
<td>□ 6.0-9.9 g/dL 18% - 27%</td>
</tr>
<tr>
<td></td>
<td>Sometimes, with significant risk factors</td>
</tr>
<tr>
<td></td>
<td>□ &gt; 9 g/dL 27%</td>
</tr>
<tr>
<td></td>
<td>Almost never, only with VERY HIGH risk factors</td>
</tr>
<tr>
<td>DIAGNOSIS - CHECK ALL THAT APPLY</td>
<td></td>
</tr>
<tr>
<td>□ Acute anemia due to acute blood loss (of &gt;750 mL) or 15% blood vol. with symptoms of hypovolemia</td>
<td></td>
</tr>
<tr>
<td>□ Acute anemia of unspecified origin (e.g. fall in HCT &gt; 6 points within 24 hours)</td>
<td></td>
</tr>
<tr>
<td>□ Anemia due to chronic disease (CIRCLE ONE)</td>
<td></td>
</tr>
<tr>
<td>□ Hypovolemic anemia</td>
<td></td>
</tr>
<tr>
<td>□ Anemia due to hemolysis</td>
<td></td>
</tr>
<tr>
<td>□ Other</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>ORDER: Type &amp; Cross-Match</th>
<th>PRBC's (Leukodepleted) units</th>
<th>Irradiated PRBC's</th>
</tr>
</thead>
<tbody>
<tr>
<td>(order units for entire treatment course)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PLATELETS</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>DIAGNOSIS - CHECK ONE</td>
<td></td>
</tr>
<tr>
<td>□ Platelet count: 10,000/µL or failure to produce</td>
<td></td>
</tr>
<tr>
<td>□ Check one: Unspecified Thrombocytopenia or</td>
<td>Specify Type</td>
</tr>
<tr>
<td>□ Platelet count: 30,000/µL or signs of hemorrhagic diathesis</td>
<td></td>
</tr>
<tr>
<td>□ Platelet count: 10,000/µL or acute hemorrhage or invasive procedure (recent or in progress)</td>
<td></td>
</tr>
<tr>
<td>□ Platelet dysfunction as documented by: (specify)</td>
<td></td>
</tr>
<tr>
<td>□ Special circumstances: (specify)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>ORDER: Platelets (Plateletpheresis)</th>
<th>units</th>
</tr>
</thead>
<tbody>
<tr>
<td>(NOT stocked at BMH - minimum of 8 hours to receive)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PLASMA</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>DIAGNOSIS - CHECK ONE</td>
<td></td>
</tr>
<tr>
<td>□ Emergent reversal of Warfarin effect or Emergent correction of elevated INR</td>
<td></td>
</tr>
<tr>
<td>□ Abnormal coagulation studies and significant hemorrhage</td>
<td></td>
</tr>
<tr>
<td>□ Other: (specify)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>ORDER: Fresh Frozen Plasma (FFP)</th>
<th>units</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CRYOPRECIPITATE</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>DIAGNOSIS - CHECK ONE</td>
<td></td>
</tr>
<tr>
<td>□ Fibrinogen: 100 mg/dl</td>
<td></td>
</tr>
<tr>
<td>□ Fibrinogen: 100 mg/dl - with active hemorrhage</td>
<td></td>
</tr>
<tr>
<td>□ Other: (specify)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>ORDER: units for entire course of treatment</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Cryoprecipitate: # units to be given</td>
<td>(NOT stocked at BMH - takes a minimum of 8 hours to receive)</td>
</tr>
<tr>
<td>Cryoprecipitate: Single unit (20 mL)</td>
<td></td>
</tr>
<tr>
<td>Cryoprecipitate: Pooled (5 units)</td>
<td></td>
</tr>
<tr>
<td>Other: (specify)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>MD Signature</th>
<th>MD Date</th>
<th>Date</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orders noted</td>
<td>RN/LPN</td>
<td>Date</td>
<td>Time</td>
</tr>
<tr>
<td>24 hr chart check</td>
<td>RN/LPN</td>
<td>Date</td>
<td>Time</td>
</tr>
<tr>
<td>Med filled to pharmacy</td>
<td>Initials</td>
<td>Date</td>
<td>Time</td>
</tr>
</tbody>
</table>

Laboratory Fax: 802-257-3865

Revision date: 12/19/91
BB - PATIENT SAMPLE COLLECTION

Blood samples submitted to the Blood Bank for testing must be properly labeled or they will not be accepted. Type and Screen and Crossmatch samples must be labeled with a Securline Blood Band at the time of collection.

The preferred specimen type is the EDTA (pink) tube. The individual collecting the sample must positively identify the patient and before leaving the bedside **MUST label the blood sample tubes with:**

1) Patient's full name (no initials or nicknames)
2) Medical Record number, social security number or date of birth
3) Date of collection
4) Time of collection
5) INITIALS of individual collecting the blood sample.

Please bear in mind that the majority of Fatal Transfusion Reactions are not due to incorrectly matched blood or immune antibodies, but rather are due to **CLERICAL ERRORS, ESPECIALLY ERRORS OF IDENTIFICATION.**
BB – PLATELETS

Platelets are transfused as a platelet pheresis (equivalent to 6-8 platelet concentrates).

Platelets should be ordered by 11:00 am for transfusion the same day Monday through Friday. If platelets are ordered by 11:00 am, platelets will be ready for transfusion by 4:30 pm.

Call the Blood Bank (ext. 8311) to order platelets.

Platelets are stored at room temperature and must be continually rotated on a rotator until infusion. Platelets should be infused as quickly as possible after they have been released from the Blood Bank and must not be stored prior to transfusion.
BB - POLICY FOR THE RETURN OF ISSUED UNITS OF BLOOD

If blood is issued for transfusion and then a change in the patient’s clinical status or other difficulty necessitates a delay in the transfusion, the following must be adhered to:

1) Return the blood to the Blood Bank as soon as possible, but always before 30 minutes has elapsed since issuance. Blood stored at 1°C to 6°C warms to 10°C in about 30 minutes at room temperature. Therefore, the blood transfusion must be either started or returned to the Blood Bank within 30 minutes from the time the unit was issued.

2) The entrance ports to the blood container must not have been penetrated or entered in any way so that sterility can be assured.

3) If the above conditions have been met, the blood may be brought back to the Blood Bank and be re-issued again when transfusion becomes possible.
BB - RELEASE OF CROSSMATCH BLOOD

Blood that is crossmatched will be held for 72 hours and then released.

One person may pick up blood for ONLY one patient at a time. Only one unit of blood will be issued per patient except in an emergency.
BB - REQUESTS

Requests for Blood Bank testing are made in the Hospital Information System through Order Entry, or by completing the Laboratory Requisition Form. The patient should have a Complete Blood Count done by this laboratory within 24 hours before transfusion.

Pre-Admission Surgical Patients

The Type and Screen or Crossmatch procedures are always performed on the day of surgery. It is important that the blood sample used for compatibility testing represents the patient’s current immunological status. Recent transfusion or pregnancy may evoke or stimulate production of unexpected antibodies.

The Blood Bank cannot accept testing results for Type and Screen or Crossmatch procedures performed at testing labs other than the BMH Bank since the samples would not be available for crossmatching if required.
Type-specific blood (blood of the same group and Rh as the patient) is generally issued for transfusion. However, if the required group and Rh are not available, ABO compatible packed red cells may be utilized. Group A, Group B, Group O or Group AB packed red cells may be transfused to an AB recipient. Group O recipients MUST receive Group O blood. Group O packed cells may be transfused to a recipient regardless of the ABO type.

An Rh positive recipient may be transfused with Rh positive or Rh negative blood. An Rh negative recipient should NOT receive Rh positive blood except in an emergency situation with Pathologist approval and only if the patient does not have anti-D.

**Informed Consent**
All patients undergoing non-emergent transfusions must be informed of the risks and benefits of blood and blood components and consent to their use. The physician should discuss the possibility of blood transfusion with the patient, the risk and benefits of transfusion, the methods whereby blood transfusion may be avoided or minimized, the positive and negative aspects of receiving homologous blood, and pre-donating and receiving autologous blood. The informed consent form documents that this discussion has taken place and must be signed by the patient.
BB - TRANSFUSION REACTIONS

Any adverse symptoms or physical signs occurring during transfusion of blood or its components should be considered potentially a part of a life-threatening reaction.

The individual hanging the blood should take the following actions immediately:

1) **STOP** the transfusion to limit the amount of blood infused.

2) Keep the intravenous line open with the infusion of normal saline.

3) **CHECK** all labels, forms and patient identification to determine if the right patient received the correct blood or component.

4) Report the suspected transfusion reaction to Blood Bank personnel immediately.

5) Complete the information on the top part of the Transfusion Reaction form and send the form to the Blood Bank as soon as possible along with the discontinued unit.

6) Hives are considered a transfusion reaction and the transfusion should be stopped if the patient develops hives.
BB - URGENT ORDER FOR BLOOD

An emergency order for crossmatched blood takes approximately 1 hour to complete if the patient does not have an antibody and if a type and screen needs to be performed. This usually requires a new blood sample to be drawn.

If a type and screen has already been performed within the last 72 hours, it takes approximately 30 minutes to crossmatch up to four units of blood.
CLINICAL CHEMISTRY

Routine and STAT Clinical Chemistry tests are performed in the main laboratory. The telephone extension is 8311. Technical personnel provide twenty-four hour coverage. This section of the Laboratory operates 24-hours a day, 7 days a week. The types of tests performed are shown below:

TESTING SCHEDULE

<table>
<thead>
<tr>
<th>Monday</th>
<th>Tuesday</th>
<th>Wednesday</th>
<th>Thursday</th>
<th>Friday</th>
<th>Sat/Sun</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immunology</td>
<td>Immunology</td>
<td>Immunology</td>
<td>Immunology</td>
<td>Immunology</td>
<td>Immunology</td>
</tr>
<tr>
<td>Therapeutic Drug</td>
<td>Therapeutic Drug</td>
<td>Therapeutic Drug</td>
<td>Therapeutic Drug</td>
<td>Therapeutic Drug</td>
<td>Therapeutic Drug</td>
</tr>
<tr>
<td>Drugs of Abuse</td>
<td>Drugs of Abuse</td>
<td>Drugs of Abuse</td>
<td>Drugs of Abuse</td>
<td>Drugs of Abuse</td>
<td>Drugs of Abuse</td>
</tr>
</tbody>
</table>

Cobas 6000: Chemistry Profiles: Basic Metabolic, Comprehensive Metabolic, Electrolytes, Hepatic and Lipid. Acetaminophen, Alanine Aminotransferase (ALT), Albumin, Alkaline Phosphatase, Ammonia, Amylase, Aspartate Aminotransferase (AST), Calculated LDL, Carbamazepine (Tegretol), Carbon Dioxide, Chloride, Cholesterol, Cortisol, Creatine Kinase, Creatinine, C-Reactive Protein (CRP), C-Reactive Protein - high sensitivity, Digoxin, Direct Bilirubin, Estradiol, Ethanol, Ferritin, Free T3, Free T4, Folate, Follicle Stimulating Hormone (FSH), Gamma Glutamate Transferrin (GGT), Gentamicin, Glucose, HDL-Cholesterol, Hemoglobin A1c, Hormone Chorionic Gonadotropin, quantitative (β-hCG), Iron, Lactate Dehydrogenase (LDH), Lactic acid, Lipase, Lithium, Luteinizing Hormone (LH), Magnesium, Parathyroid Hormone Intact (PTH Intact), Phenytin (Dilantin), Phosphorus, Potassium, Pro BNP, Prostate Specific Antigen (PSA), Protein, Salicylate, Sodium, Theophylline, Thyroid Stimulating Hormone (TSH), Tobramycin, Total T3, Total T4, Total Bilirubin, Total Iron Binding Capacity (TIBC), Triglycerides, Troponin-T, Urea Nitrogen, Uric Acid, Valproic Acid, Vancomycin, Vitamin B12, Vitamin D. CSF: Protein and Glucose.

MEDTOX: Amphetamines, Barbiturates, Buprenorphine, Benzodiazepine, Cocaine, Methadone, Methamphetamine, Opiates, Oxycodone, Phencyclidine, Propoxyphene, Tetrahydrocannabinol, Tricyclics.

Manual Testing: STAT HIV (for needlesticks and prenatal Mothers with unknown HIV status), Beta hydroxybutyrate, Fluid pH except Pleural Fluid which is performed by the Respiratory Therapy Department.
HEMATOLOGY

The Routine Hematology Laboratory performs all blood counts and coagulation testing. Additional analyses are body fluid cell counts and bone marrow preparations. This section operates 24-hours a day, with many analyses available on a STAT basis 24 hours a day.

Bone marrow biopsies are provided in conjunction with the Histology Section. Consultation on peripheral and marrow smears is available with the Pathologist.

Special stains may be considered following consultation with the Anatomic Pathology Department (Ext. 8311).

Specific Requirements: Body Fluids Other than Cerebrospinal Fluid

Body fluids other than cerebrospinal fluid should be collected in an EDTA anticoagulated tube for cell counts to prevent clotting of the specimen.
MICROBIOLOGY SECTION

Regular Microbiology personnel are on duty from 0700 to 1530 on weekdays and 0700 to 1500 on weekends and holidays. Twenty-four hour coverage is provided for emergency Gram stains, rapid group A strep testing on throat swabs, rapid antigen testing for influenza A & B, Legionella and RSV and specimen planting. Gram stains from positive blood cultures are read and reported on a 24/7 basis.

Specimen Collection

Most microbiology specimens should be collected by the physician or by the patient at home (stool & sputum samples). The laboratory does not have sufficient privacy for collections that require the patient to disrobe. Wound, drainage and skin scraping samples are best collected by the trained physician to ensure testing of appropriate material. The laboratory does not collect naso-pharyngeal swabs or washings required for pertussis, RSV testing or influenza testing.

The laboratory will collect blood cultures and clean catch urine for culture.

Generally, all specimens must be received in Microbiology within 1 to 2 hour(s) of collection. Swabs and other material refrigerated for up to 24 hours will be accepted. Special transport media for anaerobes and viral culture are available. Stool for parasite examination, white blood cells or culture must be placed in fixation within half an hour of collection. Refer to specific test pages in the Test Menu for individual guidelines.

Anaerobic Cultures

The Microbiology Laboratory processes specimens for isolation and identification of anaerobic bacteria. Specimens for anaerobic culture must be submitted in appropriate anaerobic transport media (available in Microbiology) and should be accompanied by a specimen for aerobic culture from the same site. Specimens from non-sterile sites having anaerobic bacteria as a component of the normal flora are generally not acceptable for anaerobic culture. The processing of such specimens will be considered on a case-by-case basis.

Examples of such specimens are: Throat swab, sputum or bronchoscopic specimens contaminated with upper respiratory secretions, feces or rectal swabs, urine, vaginal or cervical swab, material from abdominal wounds contaminated with upper respiratory or GI tract secretions.
MICRO - ANTIMICROBIAL SUSCEPTIBILITY TESTING

Antimicrobial susceptibility testing is performed on isolated pathogens by specific site in accordance with national laboratory standards. All culture requests are understood to be “C&S” requests. (The determination to perform an antimicrobial susceptibility test on any given isolate is made in the Microbiology Laboratory, based upon identification of the isolate and the source of the culture.) Requests for additional susceptibility testing should be made by the clinician directly to the Microbiology Laboratory. We will attempt to provide additional test results as technically possible, including sending samples to another laboratory as needed. Consultation with Infectious Disease or our Pathologists may be suggested.

Antimicrobial susceptibility testing is performed routinely using an automated system. Interpretation of these values is based upon achievable drug levels in body fluids and tissues. This information can be found in various physician handbooks and in pharmaceutical literature. Organisms, which are not suitable for automated MIC tests, will be tested by the Kirby-Bauer disc diffusion method. This method provides the S, I, R category calls only.

MICRO - BLOOD CULTURES

All routine blood cultures are screened for aerobic and anaerobic organisms. All organisms isolated from blood cultures are identified. Antimicrobial sensitivity tests are performed on most aerobic isolates. Certain organisms, when recovered, are suggestive of contamination from skin. These include coagulase negative Staph sp., diphtheroids, Propionibacterium sp., and Bacillus sp. However, clinical circumstances must be considered in evaluating the significance of any blood isolate.
MICRO - GENERAL BACTERIOLOGY

This section receives and processes specimens for routine culture. The specimens are accessioned, inoculated and incubated. Appropriate transfers are made to isolate and identify human pathogenic bacteria, and perform appropriate susceptibility testing. The sections below give a brief description of the major sub-areas of bacteriology and the other areas that comprise Microbiology.
MICRO - MISCELLANEOUS CULTURES

Specimens from normally sterile sites such as C.S.F., bone marrow, surgical specimens, joint fluids, pleural and peritoneal fluids, etc., are cultured for aerobic pathogenic organisms. Anaerobic pathogens may be detected by routine cultures; however, be aware of the special requirements for the isolation of anaerobes (see below). If an anaerobic infection is suspected, a specimen should be submitted for anaerobic culture.

Bone marrow, eye swabs, joint fluids and spinal fluids are routinely screened for fast growing aerobic pathogenic bacteria, including *Haemophilus* sp. and pathogenic *Neisseria*.

Genital tract specimens are routinely cultured to isolate *Neisseria gonorrhoeae* (when properly submitted) and aerobic pathogenic organisms. However, if *Haemophilus ducreyi*, or *Gardnerella vaginalis* is suspected, a special request should be made. A graded gram stain or wet prep is appropriate for evaluation of vaginosis as opposed to vaginal culture.

Specimens from wounds, abscesses, incisions and pus are screened for non-fastidious fast-growing aerobic organisms. If an anaerobe is suspected, a specimen must be properly submitted. See the specific anaerobic culture listings for details.
MICRO - MYCOBACTERIOLOGY (ACID FAST, TB)

Mycobacteriology deals with detection, isolation and identification of acid fast bacilli (AFB). Mycobacteria from clinical specimens and includes both smear and culture procedures. Specimens in Mycobacteriology commonly fall into one of six categories. These are:

1) Respiratory, including sputum, bronchial washings and brushings, and tracheal aspirates.
2) Urine.
3) "Sterile" pus.
4) Sterile body fluids, including blood.
5) Biopsied tissue specimens.
6) Stool / feces.

Category 1, 2 and 6 specimens are decontaminated with an alkali solution prior to inoculation on selective slants and liquid growth media. Category 3-5 specimens are incubated directly on a non-selective Mycobacterial culture slant. The slants are routinely incubated for 8 weeks before a final negative result is reported.

Acid fast smears (either concentrated or direct) are reported with 30 hours of receipt of sample in Microbiology. Culture updates are reported weekly after an initial 3 weeks of incubation.

*Mycobacterium tuberculosis* and other acid fast isolates are sent to the Vermont Department of Health for culture and susceptibility testing. Urine for AFB does not include a smear. Stool for AFB is not accepted on Fridays. Blood cultures for AFB are collected in yellow isolator tubes and sent to the University of Vermont Medical Center.
MICRO - MYCOLOGY

This area deals with the detection, isolation and identification of fungi from clinical specimens, and includes various smears and microscopic procedures for the direct detection of fungi within clinical material, as well as cultures. Specimens for Mycology commonly fall into one of several categories.

These are:
  1) Superficial scrapings and clippings including hair, nails, skin and mucous membranes.
  2) Respiratory: sputum, tracheal aspirates, bronchial washings, and brushings.
  3) Sterile body fluids.
  4) Biopsied tissues.

Category 1 and 2 specimens are inoculated onto both selective and non-selective fungal media. Category 3 and 4 specimens are inoculated onto a non-selective fungal medium. Fungal cultures are routinely incubated for 4 weeks before a final negative result is reported. KOH on category 1 specimens are reported with 30 hours. Culture updates are sent weekly. Yeast isolates are identified in the Microbiology Laboratory as either presumptive Candida albicans or Yeast not Candida albicans. Further identification or susceptibilities testing can be requested. The testing will be sent to a reference laboratory. Mold isolates are sent to a reference laboratory for identification.
MICRO - RESPIRATORY CULTURES

Nose, naso-pharyngeal swabs, throat, sputum, bronchial and tracheal aspirations are considered respiratory tract specimens. All specimens from these sites will be screened for fast-growing aerobic pathogenic organisms and certain fastidious isolates of possible clinical significance. Screening cultures and antigen detection for group A beta Strep are also available upon request.

Examples of organisms not isolated by routine culture are: Neisseria gonorrhoeae, Corynebacterium diphtheriae and Legionella pneumophila. For information on culture of non-routine organisms, see Culture Test Listing or call Laboratory if not included.

Throat specimens are to be collected so as to avoid contamination with organisms from mouth, tongue or dentures. Throat specimens designated for group A Strep will be screened for that organism only. Throat specimens for Neisseria gonorrhoeae will be screened for that pathogen only and must be plated at the time of collection.
MICRO - SPECIAL SUSCEPTIBILITY TESTS

Tests falling into this category must be specifically requested within 72 hours of submission of the clinical specimen. Blood culture isolates are an exception to this rule, being retained for 4 weeks or longer on request. Special susceptibility tests include the following:

1) Testing an isolate against antimicrobials not included in our routine panels. The Kirby-Bauer (disc method) will be utilized in this case, yielding category (not MIC) results. An additional charge will be levied in such cases. In most cases, MIC determinations for antimicrobial not included in our regular panels will have to be submitted to a reference laboratory.

2) Testing of a patient’s serum for bacteriocidal level against a clinical isolate. Specimens for such tests will be submitted to a reference laboratory.
MICRO - SPUTUM SPECIMENS

Instruct the patient to remove dentures, rinse mouth and gargle with an antiseptic mouthwash, cough deeply and expectorate into a sterile container. Cap tightly and submit to Laboratory.

All expectorated sputum specimens will be subjected to macroscopic and microscopic evaluation prior to accepting the specimen for culture. The presence of foreign bodies or ≥ 25 squamous epithelial cells per low per field will be considered grounds for rejection of the specimen based upon quality. Specimens obtained by trans-tracheal or bronchial aspiration will not be subjected to screening. Use of these collection techniques must be clearly indicated on the requisition form.
MICRO - SUBMISSION OF SPECIMENS

TECHNIQUE AND AVAILABILITY:
Specimens for Microbiology must be collected using aseptic techniques. Contamination with extraneous normal flora from the patient’s skin or with environmental organisms leads to confusing and erroneous results. Specimens must be transported to the Laboratory in sterile, leak-proof containers. We follow the Laboratory policy for acceptance or rejection of specimens. Please consult the test listing section for timing requirements for submission following collection. All specimens should be brought to the Main Laboratory or Routine specimens may be dropped off at Richards Building during the hours of 7 to 6 pm on Weekdays and at 8 to 12 pm on Saturdays. Laboratory personnel will accession tests as ordered and give to the Microbiology section.

REFLEX TESTING:
A throat culture screen for group A strep will be performed whenever an ordered group A strep antigen screen has tested negative. This reflex testing is in compliance with manufacturer’s recommendations. Because of this requirement, we ask that 2 throat swabs be collected and submitted with each group A strep antigen test request.
MICRO - URINE CULTURES

Specimens for routine urine culture (voided and catheterized) can be submitted to the Main Laboratory at any time. Urine specimens will be cultured for the aerobic, rapid growing organisms generally involved in urinary tract infections.

Non-catheterized urine samples must be obtained by specific “clean catch” method to avoid contamination by skin, fecal or vaginal organisms.

All urine samples are cultured quantitatively and a colony count is reported. Decisions for susceptibility testing are made based on colony count, purity of the organisms found and method of collection.
MICRO - VIRAL CULTURES

Specimens for viral testing must be collected and transported in special holding media, and in some cases, with rigid temperature requirements. Therefore, the laboratory should be notified in advance of any such collections so that proper arrangements can be made. Antigen detection procedures for Respiratory Syncytial Virus and Influenza A & B are available in the Microbiology Laboratory.

Specimens for viral isolation are referred to a reference laboratory.
SEROLOGY

**Manual Testing:** STAT HIV (for needlesticks and prenatal Mothers with unknown HIV status), Rheumatoid Arthritis Screen, Mononucleosis Screening, Serum and Urine hCG testing at all times and Syphilis screen (RPR) on Mondays.
URINALYSIS

The Urinalysis Section Laboratory performs routine urinalysis.

SIEMENS STATUS: Qualitative hCG.

Manual Testing: STAT HIV (for needlesticks and prenatal Mothers with unknown HIV status), Beta hydroxybutyrate, specialized urine tests as well as occult blood analysis of feces and gastric contents.
SPECIMENS REFERRED TO REFERENCE LABORATORIES

Cytopathology

If there is a delay in delivery to the Laboratory, fixed specimens should be refrigerated.

SUPPLIES:

The following supplies are available to physicians' offices from the Laboratory:

- Instructions to the Patient for urine/sputum collection
- Sterile containers for urine and sputum.
- Requisitions
- Biohazard specimen bags
- Thin Prep collection vials with spatula and Endocervical brush
- Endocervical brooms
PATHOLOGY SPECIMENS

Except for frozen sections, all pathology specimens are processed by the Dartmouth-Hitchcock network.

All specimens must be submitted to the BMH Main Laboratory accompanied by the appropriate completed requisition, including relevant history and pre-operative diagnosis. Responsibility for providing all required information rests with the clinician requesting the consultation. Please consult the BMH Specimen Collection Guide or Test Menu for appropriate fixatives and transport media.

Requisitions that may be used for the various types of pathology and cytopathology specimens are shown in the pictures on the following pages.
REQUISITION FOR ACQUIRED DISORDERS

MOLECULAR DIAGNOSTICS, CYTOGENETICS AND FLOW CYTOMETRY TESTING
Acquired Disorders

Dartmouth-Hitchcock
A CULTURE OF CARING

To: Department of Pathology & Laboratory Medicine
One Medical Center Drive
Lebanon, NH 03756-0001
Tel: (603) 650-2500, Fax: (603) 650-4646

Sent from: (REQUIRED)

Required Billing Information- Please check one box indicating who D-H is to bill

☐ Client bill to: ____________________________ (name of facility)

OR

☐ Patient bill

* Patient Billing: If D-H is to bill the patient or patient’s insurance directly, a SSN ________ ________ ________ ________ and detailed insurance information is REQUIRED (a face sheet will do)

REQUIRED INFORMATION - All Fields Are Required

Patient Name: ____________________________ DOB: __________ MRN/Account: __________

Gender (circle one): Male / Female

Status (circle one): Inpatient / Outpatient

Sample Type (circle one): Bone Marrow / Blood / Other (specify)

Date and Time Collected: ____________________________ Ordering Providers signature: ____________________________

Diagnosis: ____________________________ Status (circle one): Diagnostic / Staging / Prognosis / Remission

ICD-10 Codes: ____________________________

Tests (check all that apply):

Cytogenetics

☐ Chromosome Analysis for Acquired Disorder - Blood OR Bone Marrow in SODIUM HEPAIRIN tube at room temperature

☐ CLL panel by FISH - Blood OR Bone Marrow in SODIUM HEPAIRIN tube at room temperature

☐ FISH (specify): Blood OR Bone Marrow in SODIUM HEPAIRIN tube at room temperature

☐ BCR/ABL1 Testing by FISH (detected p190 and p210) - call 603-650-7761 with questions

Molecular Pathology

☐ BCR/ABL1 p210 Testing by RT PCR - call 603-350-8257 with questions

☐ JAK2 Mutation for myeloproliferative disorder (MPD) or polycythemia vera (PV)

☐ FLT3 or AML N-karyotypic collection tubes - NO HEPARIN TUBES can be accepted

☐ IGH Gene Rearrangement (PCR, EDTA or ACD are acceptable collection tubes - NO HEPARIN TUBES can be accepted

☐ TCR Gene Rearrangement (PCR)

☐ FLT3 or AML N-karyotypic collection tubes - NO HEPARIN TUBES can be accepted

☐ BCL2 (PCR, t(14;18):

☐ EDTA or ACD are acceptable collection tubes - NO HEPARIN TUBES can be accepted

How Cytometry

☐ Flow Cytometry immunophenotyping for Chronic lymphocytic leukemia disease (specify):

Blood OR Bone Marrow in an EDTA tube at room temperature

Has a helpful PHENOTYPIC CLINICAL with constant platelets

☐ Flow Cytometry immunophenotyping for Acute lymphocytic/myelogenous leukemia disease (specify):

Blood OR Bone Marrow in an EDTA tube at room temperature
REQUISITION FOR INHERITED CONGENITAL DISORDERS

Required Billing Information - Please check one box indicating who D-H is to bill:

☐ Client bill to: ___________________________ (name of facility)

☑ Patient bill

* Patient Billing: If D-H is to bill the patient or patient’s insurance directly, a SSN: ___________________________

and detailed insurance information is REQUIRED (a face sheet will do).

REQUIRED INFORMATION – All Fields Are Required

Patient Name: ___________________________ DOB: ___________ MRN/Account: ___________________________

Gender (circle one): Male / Female

Status (circle one): Inpatient / Outpatient

Sample Type (circle one): Blood / Tissue (specify source): ___________________________

Date and Time Collected: ___________________________ Ordering Provider’s Signature: ___________________________

*Diagnosis: ___________________________

ICD-10 code(s): ___________________________

Test(s) Requested (check all that apply):

☐ Chromosome Analysis / Karyotype

☐ FISH (specify):

Blood: 5-15 ml whole blood in KEDBA or EDTA tube at room temperature

☐ Chromosome Microarray Analysis

Blood: 5-15 ml whole blood in EDTA tube at room temperature

☐ Fragile X

Blood: 5-15 ml whole blood in an ACD tube at room temperature

☐ Factor V Leiden

Blood: 5-15 ml whole blood in an EDTA, ACD, or Citrate tube at room temperature

☐ Hcm analysis

Blood: 5-15 ml whole blood in an EDTA or ACD tube at room temperature

☐ Prothrombin

Blood: 5-15 ml whole blood in an EDTA, ACD, or Citrate tube at room temperature

☐ MTHFR (methylene tetrahydrofolate reductase mutation)

Blood: 5-15 ml whole blood in an EDTA, ACD, or Citrate tube at room temperature

☐ Cystic Fibrosis / DNA (specify): Diagnose Cystic Fibrosis consent form – call 603-650-3269 for info

Blood: 5-15 ml whole blood in an EDTA or ACD tube at room temperature

☐ UST/USI – UST Polymorphisms

Blood: 5-15 ml whole blood in an EDTA or ACD tube at room temperature

☐ DNA Archive

Blood: 5-15 ml whole blood in an EDTA or ACD tube at room temperature

☐ SRY by PCR

Blood: 5-15 ml whole blood in an EDTA or ACD tube at room temperature
### Cytopathology GYN

**Referring Identifier:**
- [ ] Patient Full Legal Name (First and Last Name)
- [ ] DH MRN

**Copy to:**
- [ ] Date of Birth
- [ ] Sex
  - [ ] Male
  - [ ] Female
- [ ] Patient Type
  - [ ] InPt
  - [ ] OutPt
- [ ] Location
- [ ] Skilled Nursing Facility (SNF)?
  - [ ] Yes
  - [ ] No

**Ordering Provider (First and Last Name):**
- [ ] Ordering Provider Signature
- [ ] Collector Signature
- [ ] Collection Date and Time

**Billing provider (First and Last Name), if different**

### INFORMATION REQUIRED FOR PROCESSING GYN SPECIMENS:
Choose selections from each of the four categories:

1. **Circle ICD-10 Code, Choose ONE:**
   - [ ] Z12.4 Encounter for screening for malignant neoplasm of cervix
   - [ ] Z91.89 Hx. Clin. High Risk
   - [ ] Z85.40 Hx. other malignancy, female genital tract
   - [ ] N80.9 Endometriosis
   - [ ] N91.2 Amenorrhea
   - [ ] N96.9 Meno or postmenop dis.
   - [ ] N73.9 PID
   - [ ] R87.620 Vaginal Pap with ASCUS
   - [ ] N96 Non-inflam. disorder
   - [ ] R87.621 Vaginal Pap with ASCH
   - [ ] N92.6 Menstrual disorder
   - [ ] R87.622 Vaginal Pap with LGSIL
   - [ ] N24.79 Encounter for screening for malignant neoplasm of other genitourinary organs
   - [ ] R87.623 Vaginal Pap with HGSIL
   - [ ] Z12.69 Encounter for screening for malignant neoplasm of other sites
   - [ ] Other:

2. **HPV testing options, choose ONE based on patient’s age:**
   - [ ] Pap Testing ONLY. Do not perform High Risk HPV testing unless notified by provider*
   - [ ] Perform High Risk HPV testing ONLY
   - [ ] Perform High Risk HPV if Pap test result is ASCUS (ASCUS triage)
   - [ ] Perform Concurrent Pap and High Risk HPV testing (Primary HPV screening)
   - [ ] *Note: Sample will be discarded 30 days from collection date. Call 603-650-2200 to order HPV testing

3. **Specimen Information/Relevant Clinical Information**
   - [ ] Liquid Based Pap
   - [ ] Vaginal
   - [ ] Cervical
   - [ ] Endocervical
   - [ ] Collection Date:
   - [ ] LMP:
   - [ ] Postmenopausal
   - [ ] Yes
   - [ ] No
   - [ ] Currently on Gyn hormones (estrogen/progestosterone)
   - [ ] Yes
   - [ ] No
   - [ ] Hysterectomy?
   - [ ] Total
   - [ ] Supracervical
   - [ ] Yes
   - [ ] No
   - [ ] Pregnant
   - [ ] Yes
   - [ ] No
   - [ ] IUD
   - [ ] Yes
   - [ ] No
   - [ ] Pelvic Radiation
   - [ ] Yes
   - [ ] No
   - [ ] Prior GYN therapy (cone bx, cautery, cryotherapy, surgery)
   - [ ] Yes
   - [ ] No
   - [ ] Previous abnormal Pap or cervical biopsy
   - [ ] Yes
   - [ ] No
   - [ ] History of HPV vaccination?
   - [ ] Yes
   - [ ] No
   - [ ] History of smoking?
   - [ ] Yes
   - [ ] No
   - [ ] Previous DES exposure
   - [ ] Vaginal Smear

4. **Pertinent Clinical Data, Clinical Impression and Significant Therapy:**
   - [ ] Pap after ASCUS
   - [ ] Pap after CIN HPV on bx
   - [ ] Pap after cryo, Leep, CCK for CIN

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Phone: 603-650-2200 — Fax: 603-650-8151 — [http://www.dartmouth-hitchcock.org](http://www.dartmouth-hitchcock.org) — CLIA ID# 30D0652833 H-2536

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BMH LAB GUIDE

GENERAL INFORMATION

November 2018
# Requisition for Cytopathology Non-Gyn Specimens

<table>
<thead>
<tr>
<th>Fluids:</th>
<th>Gastrointestinal Brushings (endoscopies):</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pleural Site (L,R)</td>
<td>Gastric Site (L,R)</td>
</tr>
<tr>
<td>Peritoneal Site (L,R)</td>
<td>Esophageal Site (cm)</td>
</tr>
<tr>
<td>Pericardial</td>
<td>Colon Site (cm)</td>
</tr>
<tr>
<td>Bronchial Wash Site (L,R,M)</td>
<td></td>
</tr>
<tr>
<td>Bronchial Brush Site (L,R,M)</td>
<td></td>
</tr>
<tr>
<td>BAL Site (L,R,M)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Urines:</th>
<th>Miscellaneous:</th>
<th>Fine Needle Aspiration:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Voided</td>
<td>Other Site</td>
<td>Site – REQUIRED INFORMATION</td>
</tr>
<tr>
<td>Catheterized</td>
<td>Scrapings Site</td>
<td></td>
</tr>
<tr>
<td>Cystoscopic Site</td>
<td>Washings Site</td>
<td></td>
</tr>
</tbody>
</table>

### Clinical History:

### Clinical Impression:

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F-312 9/15
REQUISITION FOR SURGICAL, DERMSPATH SPECIMENS

<table>
<thead>
<tr>
<th>Surgical Pathology/Dermatopathology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Referring Identifier:</td>
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<tr>
<td>Patient Full Legal Name (First and Last Name)</td>
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<td>DH MRN</td>
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<td>Collector Signature</td>
</tr>
<tr>
<td>Billing provider (First and Last Name), if different</td>
</tr>
</tbody>
</table>

**Specimen Type:**
(Note: Multiple biopsies from different regions should be submitted in separate bottles. Multiple pieces from one site may be submitted in one bottle.)

- **Surgical Pathology:** Biopsy □ Excision □ Resection
- **Dermatopathology:** Shave □ ED&C □ Excision □ Re-excision

If Breast biopsy – please provide time of excision and time into fixative (Ischemic Time)

Specimen(s) Submitted:

Clinical Information:

Clinical Diagnosis:

Phone: 603-650-2200 — Fax: 603-650-8151 — [Website] — CLIA ID# 30D0652833 — H-2573