# BLOOD COLLECTION

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1. Background and Rationale

The year 3 standard blood collection is much simpler than in previous years, amounting to the collection of a pediatric tube for a complete blood count, and one 10 cc serum tube for archival purposes. Any participants for whom we did not collect the buffy coats and platelets using the final cell protocol, and who did not refuse this collection, should have 2 CPT tubes drawn as well.

The Health ABC study year 3 clinic visit involves the collection of approximately 20 mL of blood (including blood for the arterialized blood gas protocol), unless the cell protocol was skipped in year 2. With the collection of blood for the cell protocol, the total volume is 36 mL. Since the study depends on the voluntary return of participants over an extended period of time, every effort must be made to make the entire procedure as easy and painless as possible both for the participants and for the Field Center personnel.

In most cases, the blood will be drawn as part of the arterialized blood gas protocol (see Chapter 2C). However, if the participant did not have cell collection in Year 2 (see Data from Prior Visits report) and is willing to have it in Year 3, the blood draws should be done separately, since the total volume is too great to perform the draw comfortably through a hand vein.

If the blood will be drawn as part of the arterialized blood gas protocol, the standard blood samples will be withdrawn with syringes after inserting the butterfly, but before flushing with heparin. Do not use a vacutainer when collecting blood from the hand veins, as this is likely to lead to vein collapse or damage and will preclude being able to draw an arterIALIZED sample. Details of the blood draw in conjunction with the arterialized blood gas protocol are given in Chapter 2C.

If a participant refuses cell collection, refuses the arterialized blood gas protocol, or two attempts were made without success to get an arterialized sample, the collection protocol outlined below should be used.
A standard informed consent has been prepared for this study. With regard to laboratory procedures, the consent statement informs study participants that there is a small risk of bruising at the spot on the arm where the blood is taken and that about three tablespoons of blood are drawn. The consent statement also informs study participants that they will be contacted if clinically significant test results are discovered.

2. Equipment and Supplies

2.1 Sample ID Labels

You will be supplied with sheets of sample ID barcode labels to use for labeling forms, draw tubes, and cryovials. A sample sheet of barcode labels can be found in Appendix 1. All labels on each sheet have the same 6-digit sample ID number (the first digit identifies the clinic: Memphis = 1, Pittsburgh = 2).

There are 12 labels containing the ID number with no extension. These labels have no barcode, and they have 1-3 lines of text indicating which specimen container they are intended for, including the stopper color and volume, if applicable. Four of these labels are to be used for pre-labeling the 4 draw tubes, with 3 extras for backup vacutainers, one for the citrate pooling tube, and four for labeling the conical centrifuge tubes used for buffy coat and platelet separation. If no CPT tubes need to be drawn (participant had cells collected during Year 2), these last 5 labels and the two CPT-tube labels can be discarded.

There are also 2 barcoded labels with the ID number, one with the words “Phlebotomy Form,” which is placed on the Phlebotomy Form (Appendix 2), and the other with the words, “Laboratory Processing Form” which is placed on the Laboratory Processing Form. This process of matching the participant-specific Health ABC Enrollment ID # (already on the form brought to the lab by the participant) to the sample-specific ID barcode is crucial to being able to use the data collected from laboratory tests.

There are also 3 barcoded labels with the same ID number and the words “BDID Form.” Use of these labels is detailed in the Lab Specimen Processing chapter. Finally, there are 12 labels intended for labeling cryovials (see Lab Specimen Processing chapter). Again, if the participant does not need a CPT draw, the two platelet, two buffy coat, and 4 citrate labels can be discarded. All of the labels relating to CPT collection and processing are grouped together at the bottom of the
sheet of labels so that they can be easily discarded or separated for the cell processor to use.

2.2 Blood Collection Trays and Tubes

Blood drawing trays are prepared in advance for the following day. Each tray is stocked with a full supply of blood drawing equipment for 6-9 participants and holds an ice bath and the individual blood collection tube rack for each participant. Several racks will also be prepared to hold various plastic tubes and vials for the final aliquots sent to the Laboratory for Clinical Biochemistry Research (LCBR) or to McKesson BioServices for archiving. The blood collection tube racks and aliquot tube racks are prelabeled from the same sheet of sample ID barcode labels.

2.2.1 Blood Collection Tray

The collection tray itself is made of hard plastic, which is unbreakable and can be easily cleaned. The tray has ten individual compartments, which are filled with the following supplies:

- Alcohol swabs
- Band-Aids
- Gauze
- Tourniquets (2)
- Vacutainer holders
- Needle/ sharps container
- Styrofoam ice bath filled ~10 min before draw
- 21G Butterfly needles with Luer adapter
- Smelling salts
- Timer/ stopwatch
- Scissors
- Adhesive tape
- Pencils/ pens
- Latex gloves

2.2.2 Blood Collection Rack: Labeling and Setup

A separate tube rack containing the necessary draw tubes is set up for each participant. They are arranged according to the priority of the draw. This rack will fit into the blood collection tray. The blood collection tubes should be prelabeled with sample ID labels. Note that the vacutainers will still be used in conjunction with the arterialized blood gas protocol, just not directly connected to the butterfly (see Chapter 2C). After withdrawal of the sample into a syringe, the blood will be transferred to the vacutainers by attaching a new needle to the syringe and piercing the stopper of the vacutainers. Thus, vacutainers can be prelabeled without regard to whether the participant will have a successful arterialized sample collection.
After the labels have been used for setting up the blood collection rack and the aliquot rack (see Lab Specimen Processing chapter), there will be 8 labels left: 3 “Backup Vacutainer” labels, 1 “Phlebotomy Form” label, 1 “Laboratory Processing Form” label, and 3 “BDID Form” labels. These can be separated into 2 mini-sheets: The “Backup Vacutainer,” “Phlebotomy Form,” and “Laboratory Processing Form” labels should be clipped to the corresponding blood collection tray. The labels for the pooling tubes, conical centrifuge tubes, and the BDID Form should be clipped to the corresponding aliquot rack.

2.2.3 Description of Blood Collection

Each draw tube is color coded to aid in handling.

Tube 1 is a 3-mL pediatric tube for the complete blood count (CBC). After drawing, the tube is mixed briefly and placed on ice.

Tube 2 and 3 are special 8 mL cell separation (CPT) tubes with blue/black stoppers. These will only be used for participants who did not have cells collected in Year 2 and who did not refuse the cell collection protocol. See Health ABC Data from Prior Visits to determine whether the participant requires these two tubes. The CPT tubes will be processed to obtain viable buffy coats for later studies of intracellular markers of aging and disease and for production of DNA for genetic analyses. Platelets will also be collected for analysis of mitochondrial enzymes and DNA. These tubes use citrate as the anticoagulant, so the plasma from these tubes is aliquoted into blue-capped cryovials. CPT tubes are supplied by LCBR and may be stored at room temperature.

Tube 4 is a 10 mL siliconized red stoppered tube. This tube contains no anticoagulant so that the blood clots to form serum. After drawing, the blood is allowed to clot at room temperature for 40-45 minutes (Maximum = 90 minutes). Cryovial caps are coded red. The serum is used for archiving.
3. Safety Issues and Exclusions

3.1 Precautions for Handling Blood Specimens

In accordance with the OSHA regulations on blood borne pathogens (see Appendix 4 for complete OSHA regulations), the LCBR recommends the following laboratory safety protocol for the field center laboratories:

- Non-permeable lab coats, latex gloves, and face shields should be used when handling any blood in any situation where splashes, spray, spatter, or droplets of blood may be generated and eye, nose, or mouth contamination can be reasonably anticipated.

- 'Universal Precautions' should be followed when handling any blood products.

- Contaminated needles and sharps shall be immediately placed in a puncture-resistant, leakproof container. Never recap or break needles.

- Hepatitis B vaccine should be offered to all unvaccinated technicians handling blood and documentation of vaccination or technicians declining to be vaccinated should be kept.

3.2 Participant Precautions and Exclusions

3.2.1 Participant Phlebotomy Questionnaire  The first question on the Phlebotomy form helps you determine whether the participant requires blood drawn for the cell collection protocol (tubes #2 and 3). The information required can be found on the Health ABC Data from Prior visits report. Next, following the questionnaire format on the Phlebotomy Form, each participant is asked whether they have a bleeding disorder before the blood is drawn (Ques. 2). If they have had any problems with excessive bleeding or bruising at a venipuncture site, use your own judgment to decide whether or not a clinic physician or nurse supervisor should be consulted.

If the participant has experienced fainting spells during phlebotomy (Ques. 3), ask the participant the frequency of fainting spells. If the participant frequently faints, again, use your own judgment to determine whether or not a consultation with the clinic physician or nurse supervisor is necessary. Provide smelling salts, basin, and
a cold cloth if needed. See section below on precautions when a participant feels faint.

Questions 4 and 5 relate to rare, but important exclusions. If a participant has had a radical mastectomy, including removal of the axillary (armpit) lymph nodes, any damage to veins on the side from which the lymph nodes were removed could result in chronic edema and clotting problems. Therefore, it is safest to use the arm from which lymph nodes were not removed. If a participant has had a bilateral radical mastectomy, it is safest not to do the blood draw at all. If they aren't sure whether their mastectomy was radical or a modified procedure, it is safest to treat it like a radical mastectomy.

Similarly, if a participant has had a graft to allow kidney dialysis, application of the tourniquet and venipuncture in the area of the graft could seriously compromise the graft. Again, it is safest to use the arm without the graft or, if the participant has had grafts in both arms, not to do the blood draw at all.

3.2.2 PRECAUTIONS WHEN A PARTICIPANT FEELS FAINT OR LOOKS FAINT FOLLOWING THE BLOOD DRAWING

- Have the participant remain in the chair; if necessary have them sit with their head between their knees.
- Provide the participant with a basin if they feel nauseated.
- Have the participant stay sitting until the color returns and they feel better.
- Place a cold wet cloth on the back of the participant's neck.
- If the participant faints, use smelling salts to revive by crushing the ampule and waving it under the participant's nose for a few seconds.
- If the participant continues to feel sick, contact a medical (nursing) staff member who will advise you on further action.

3.3 Participant Refusal of Phlebotomy

Rarely, a participant will refuse phlebotomy. Please keep a list of Health ABC Enrollment ID #s of any of these participants.
4. Participant and Exam Room Preparation

4.1 Phlebotomy Room

The blood drawing should take place in an isolated room, or participants should be separated by room dividers. The room should be equipped with all of the necessary blood drawing supplies. A separate counter or work table should be equipped with all of the materials and vials that are used in the blood handling and processing. The centrifuge, refrigerator, and freezer should be nearby.

4.2 Preparation for Phlebotomy

Preparation for phlebotomy is done in the following manner. Early morning, before any participants arrive:

- Check to make sure that the blood collection tray is properly equipped. Every item on the checklist (Appendix 5) must be ready before proceeding.
- Check that each vacutainer tube is properly labeled with sample ID labels and ordered 1-4.
- Check that the sample processing station is properly equipped (see Lab Sample Processing chapter).
- Make sure the phlebotomy room is tidy and stocked with extra smelling salts, basin, and disposable wash cloths, and that the blood mixer is functional.

Approximately 10 minutes before scheduled participant arrival:

- Fill styrofoam ice bath 3/4 full with crushed ice.

4.3 Preparation of Participants for Phlebotomy

It should be stressed that this study requires the voluntary cooperation of the participants. These people are donating both time and blood on a purely voluntary basis, with no reward other than the knowledge that they are contributing to progress in medicine. Thus, the whole experience must be made as pleasant as possible. Two to four tubes of blood of various sizes are collected, each containing
about 1-2 teaspoons (5-10 mL) of blood. Any participants who are concerned about the volume of blood should be reassured that the total amount of blood drawn is about 2-3 tablespoons, although it may look like more. The phlebotomist may also assure participants that they donate more than 12 times as much blood (450 mL) when they donate a unit of blood.

5. Detailed Procedures

5.1 Forms

An example of the Phlebotomy form is in Appendix 2. The purpose of this form is to provide a vital link between the sample ID# and the participant ID# and to facilitate the collection of plasma and serum samples from participants. The collection must be done in a rapid and efficient manner, with maximum protection for the participant. In addition, the process must facilitate the monitoring of phlebotomy and other quality assurance parameters. All forms must be completed in ink.

The Phlebotomy form has the following purposes:

1. Determine and record whether the participant requires cell collection in year 3.
2. Assure the most efficient and safest possible venipuncture for participants.
3. Allow the monitoring of the quality of the above procedures.
4. Provide information critical to the interpretation of future assay results.
5. Provide the vital link between the sample-specific barcode and the participant-specific Enrollment ID#.

The participant will arrive at the phlebotomy station with their Health ABC Enrollment ID# already filled in on their Phlebotomy and Laboratory Processing forms. The sample ID will be determined by the set of prelabeled tubes used to collect their samples. It is vital that this same sample ID be matched up with the participant’s Health ABC Enrollment ID# on both the Phlebotomy and the Laboratory Processing forms (see Lab Specimen Processing chapter). There will be a small sheet of labels clipped to the rack of vacutainers. On it is a “Phlebotomy Form” label, to be affixed to the upper right corner of the Phlebotomy Form, and a “Laboratory Processing Form” label, which should be affixed to the lower left corner of the Laboratory Processing Form. This should be done before drawing any blood, to insure that this critical task is not forgotten.
There are actually two parts to the Phlebotomy form associated with blood drawing. The first section contains questions that are important for participant safety; these questions should be asked immediately before phlebotomy and deal with any propensity to bleed or faint. The second part deals with details of the phlebotomy procedure, whether it went smoothly, how long it took, etc.

5.1.1 Return Visits for Sample Collection

Occasionally, participants may be unwilling or unable to provide a blood specimen at the time of the visit, but willing to return to the clinic for this purpose. The most important specimens are the buffy coat and platelets (if not collected during Year 2). If for some reason a participant cannot have a blood draw at all at the time of the clinic visit but is willing to return for phlebotomy, this should be strongly encouraged for the sake of the cell specimens. However, if the participant already had cell specimens collected in Year 2 and does not need to return to the clinic for any other reason, it is not necessary to have them return just for a non-cell Year 3 blood draw.

There are separate forms that must be filled out for return visits: the Return Visit Phlebotomy Form and the Return Visit Lab Processing Form (see Appendix 3). Use a new set of sample ID bar code labels. Place the Phlebotomy Form label in the Bar Code Label space on the Return Visit Phlebotomy Form. Place a Laboratory Processing Form label in the Bar Code Label space on the Return Visit Laboratory Processing Form. Use the draw tube and cryovial labels as usual. Be sure to fill out both forms with the header information including the Health ABC Enrollment ID #, Acrostic, Date Form Completed, and Staff ID #.
5.2 Phlebotomy

5.2.1 General

The venipuncture is performed with a 21 gauge butterfly needle with 12 inches of plastic tubing between the venipuncture site and the blood collection tubes. A 23 gauge needle may be used, if necessary, for a difficult draw, but this must be noted on the Phlebotomy form under “Comments on blood collection.” The butterfly has a small, thin-walled needle, which minimizes trauma to the skin and vein. The use of 12 inches of tubing allows tubes to be changed without any movement of the needle in the vein. If the participant is concerned about the venipuncture, they may be reassured to know such care is taken. The participant should be given enough time to feel comfortable both before and after the blood collection. In many cases the most memorable part of the experience for the participant will be the contact with the technician who draws the blood and their general attitude and competence.

If the participant is nervous or excited, the technician briefly describes the procedure. Sample script: “I am going to be drawing about 2-3 tablespoons of blood. This blood will be used in tests for a complete blood count and to store for later tests. We hope to be able to use the results of these tests to better understand health and disease in older people.”

5.2.2 Handling participants who are extremely apprehensive about having blood drawn

Do not under any circumstances force the participant to have blood drawn. It may help to explain to the participant that the blood drawing is designed to be as nearly painless as possible. It is sometimes best to let the participant go on with another part of the visit. It may also be helpful to have the participant relax in the blood drawing chair just so the phlebotomist can check the veins in the participant's arms, without actually drawing blood. If the participant has “good veins” the phlebotomist can reassuringly say, “Oh, you have good veins; there should be no problem.” Elderly participants are often aware of the difficulty they pose to phlebotomists and should receive extra consideration and detailed explanations as required.

5.2.3 Venipuncture Procedure

• Wear Latex gloves and a lab coat.
• Arrange draw tubes in order of draw (see below) on the table top within easy reach. Assemble butterfly apparatus and vacutainer holders, gauze, and alcohol prep prior to tourniquet application.
• Apply tourniquet.
• Examine participant's arms for the best site for venipuncture. Generally the antecubital vein is preferred, if feasible. Release tourniquet.
• Cleanse venipuncture site. Prepare area by wiping with alcohol swab in a circular motion from center to periphery. Allow area to dry.
• Reapply tourniquet and start timer. Note the start time on the Phlebotomy form. Use standard 12-hour time and include am/ pm.
• Grasp the participant's arm firmly, using your thumb to draw the skin taut. This anchors the vein. The thumb should be 1 or 2 inches below the venipuncture site.
• With the needle bevel upward, enter the vein in a smooth continuous motion.
• Make sure the participant's arm is in a flat or downward position while maintaining the tube below the site when the needle is in the vein. It may be helpful to have the participant make a fist with the opposite hand and place it under the elbow for support.
• Grasp the flange of the vacutainer holder and push the tube forward until the butt end of the needle punctures the stopper, exposing the full lumen of the needle.
• Note the blood flow into the first collection tube. If blood is flowing freely, the butterfly needle can be taped to the participant's arm for the duration of the draw. If the flow rate is very slow, the needle may not be positioned correctly.
• Remove the tourniquet at 2 minutes. Once the draw has started, do not change the position of the tube until it is withdrawn from the needle. If blood flow ceases after the tourniquet is removed, it may be reapplied for another 2 minutes. Note on the Phlebotomy form the total length of time the tourniquet was on. If a second stick is required because no blood was obtained with the first venipuncture, record the time the tourniquet was on the second arm. If a second stick is required because flow stopped or the draw was in some way interrupted, record the sum of the tourniquet times for both arms.
• Keep a constant, slight forward pressure (in the direction of the needle) on the end of the tube. This prevents release of the shutoff valve and stopping of blood flow. Do not vary pressure or reintroduce pressure after completion of the draw.
• Fill each vacutainer tube as completely as possible; i.e., until the vacuum is exhausted and blood flow ceases. If a vacutainer tube fills only partially, remove the vacutainer and attach one of your extra, backup tubes of the same type without removing the needle from the vein. Be sure to place one of the "Backup Vacutainer" labels on that tube after completing phlebotomy.
• When the blood flow ceases, remove the tube from the holder. The shutoff valve re-covers the point, stopping blood flow until the next tube is inserted.
• As each tube is removed, mix by gently inverting before placing tube on the mixer. Note: do NOT mix red top tube #4. (See section on Blood Mixing During Venipuncture below).
• Average venipuncture time is 2-4 minutes, but any difficulties may increase this time to 10 or 15 minutes. Be sure to note the time venipuncture is completed on the Phlebotomy form. Use standard 12-hour time and include am/pm. Please note that the end of draw equals the end of draw of conventional (non-acidosis) samples (either when the needle is removed from the arm if antecubital, or when the last syringe of blood that will be transferred to the serum vacutainer (tube #4) is removed from the butterfly (if combined with the hand stick).

5.2.4 Removing the Needle

• To remove the needle, lightly place clean gauze over venipuncture site. Remove the needle quickly and immediately apply pressure to the site with a gauze pad. Discard needle into puncture-proof sharps container.
• Have the participant hold the gauze pad firmly for one to two minutes to prevent a hematoma.
• Remove CBC tube (#1) from the blood mixer and place on ice.
• Remove CPT tubes (#2 and 3, if used) from the blood mixer and place in a rack at room temperature. Tube #4 should already be in the rack at room temperature.

5.2.5 Bandaging the Arm

Under normal conditions:
• Slip the gauze pad down over the site, applying mild pressure.
• Apply an adhesive or gauze bandage over the venipuncture site after making sure that blood flow has stopped.
• Tell the participant to leave the bandage on for at least 15 minutes.

If the participant continues to bleed:
• Apply pressure to the site with a gauze pad. Keep the arm elevated until the bleeding stops.
• Wrap a gauze bandage tightly around the arm over the pad.
• Tell the participant to leave the bandage on for at least 15 minutes.

5.2.6 Completing the Blood Drawing Procedure

• Dispose of needle and tubing in the appropriate biohazard needle sharps containers.
• Complete the first page of the Phlebotomy form. This includes rating the venipuncture as clean or traumatic, checking which collection tubes were filled, and writing comments about any difficulties with the phlebotomy under “Comments on Phlebotomy.”
• Clean up the venipuncture area (if necessary).
• Bring blood collection tray to the processing area with the filled vacutainer tubes and Laboratory Processing form.

5.2.7 Procedures for Difficult Draw

If a blood sample is not forthcoming, the following manipulations may be helpful.

• If there is a sucking sound, turn needle slightly or lift the holder in an effort to move the bevel away from the wall of the vein.
• If no blood appears, move needle slightly in hope of entering vein. Do not probe. If not successful, release tourniquet and remove needle. A second attempt can be made on the other arm.
• Loosen the tourniquet. It may have been applied too tightly, thereby stopping the blood flow. Reapply the tourniquet loosely. If the tourniquet is a velcro type, quickly release and press back together. Be sure, however, that the tourniquet remains on for no longer than two minutes at a time.
• DO NOT attempt a venipuncture more than twice unless a participant encourages you to do so.
• Reassure the participant that the inability to obtain a clean venipuncture is not any sign of a medical problem on their part.
• If venipuncture is unsuccessful and cell collection was required, participant should be rescheduled at a later date, preferably with a different Field Center phlebotomist.
• Document any problems with venipuncture and sample collection on the Phlebotomy form. In particular, note whether a vein other than one of the antecubital veins or a hand vein in connection with the arterialized blood gas protocol was used.

If a full blood draw is not possible, and it is unlikely that returning at a later date can be done, prioritize the blood draw: If a participant did not have their cells collected during Year 2 of the Health ABC study and did not refuse DNA collection, then collecting the cells (along with the serum for the regular draw) would take priority. If, however, the cells had already been collected during the Year 2 visit, or the participant refused DNA collection, the arterialized venous blood gas sample would be the first priority.
5.2.8 Other Possible Problems

1) Not all tubes are collected (blood flow ceases, difficult venipuncture, etc.): Always fill the collection tubes in the order specified. Make notations of difficulties on the Phlebotomy form. If the participant is willing, another attempt should be made to complete the draw. Remember to record the sum of the two tourniquet times (see section 5.2.3).

2) Collection tube does not fill: First, try another tube of the same type. Partially filled CPT tubes are not acceptable if at least half full. Partial tubes for serum are okay, but will result in a reduced number of aliquots. If a tube is not completely filled, check “No” (not filled to capacity) and explain why under Question 15 of the Phlebotomy form.

5.2.9 Priority of Tubes

A total of approximately 13-29 mL of blood will be drawn from each participant in 4 tubes. Tubes are numbered 1-4 and arranged in the rack to be drawn in the following order of priority:

1. CBC 3 mL purple top
2. CPT 8 mL blue and black top
3. CPT 8 mL blue and black top
4. Serum 10 mL red top

5.3 Blood Mixing During Venipuncture

Each tube should be treated as follows:

#1 CBC: mix gently, then place on ice. Insufficient mixing may cause platelet clumping and an inaccurate platelet count.
#2 & 3 CPT: mix gently, then place in rack at room temperature
#4 Serum: do NOT mix; place in rack at room temperature for AT LEAST 40 minutes

6. Procedures for Performing the Measurements at Home
This examination can be done on home visits. The timing of the draw should be arranged so that there is sufficient time to be sure that the participant will not faint or bleed after the examiner has left the home, but late enough in the visit so that the samples can be returned to the lab for processing within two hours after the draw.
7. Alert Values/Follow-up

Most of the blood collected in Year 3 is for archival purposes. Besides the blood gas results, which will be measured at the Field Centers (see Chapter 2C), the only analysis being done immediately will be the Complete Blood Count (CBC). The CBC will be done at a local laboratory and results reported to the field centers by mail within 14 days. Upon receipt of the report, the data need to be transcribed onto the CBC Results form (see below) and scanned into the data system. Be sure the units on the report match those on the form. If not, the results must be converted so that they are recorded on the form in the correct units.

Two copies of the CBC report are sent by the local laboratory, one for the participant's file and one to be sent to the participant. If an abnormal value (see "Immediate Alerts" in the following table) is detected, however, the laboratory notifies the field center by both phone and fax.

Platelet clumping:
Sometimes the CBC sample clumps a bit and the laboratory is unable to get an accurate platelet count, but they are able to give a qualitative reading, such as
“sufficient.” Pittsburgh’s lab does not report a platelet count, but categorizes the platelets as “normal,” “sufficient,” or “insufficient.” Memphis’ lab does report a platelet count, but adds a footnote warning that these results may not be reliable. If the platelet count reported by the Memphis lab is \(\geq 50,000\), the report warns that platelet clumping was observed and re-evaluation may be helpful. If the count reported is \(< 50,000\), the report warns that they were unable to determine an accurate platelet count and requests that you redraw the blood. It is not necessary to ask the participant to return for a redraw. In the event that the qualitative reading is “normal” or “sufficient” (Pittsburgh) or \(\geq 50,000\) with a warning (Memphis), please record –66 in the “platelets” boxes on the CBC Results form. If the qualitative reading is “insufficient” (Pittsburgh) or \(< 50,000\) with a warning (Memphis), please record –55. Be sure to include the minus (–) sign.

Mean corpuscular volume:
If the local laboratory reports mean corpuscular volume to decimal accuracy you should round to the nearest whole number. Round up if .5 or greater; round down if .4 or less (e.g., 79.8 would be rounded to 80, 79.3 would be rounded to 79).

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Reference Range for Reports</th>
<th>Immediate Alerts*</th>
</tr>
</thead>
<tbody>
<tr>
<td>White blood cells</td>
<td>See local lab reference range</td>
<td>&lt;2,000 or &gt;15,000</td>
</tr>
<tr>
<td>Red blood cells</td>
<td>See local lab reference range</td>
<td>None</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>See local lab reference range</td>
<td>None</td>
</tr>
<tr>
<td>Hematocrit</td>
<td>See local lab reference range</td>
<td>&lt;30% or &gt;50%</td>
</tr>
<tr>
<td>Platelets</td>
<td>See local lab reference range</td>
<td>&lt;100,000 or &gt;600,000</td>
</tr>
<tr>
<td>Mean corpuscular vol.</td>
<td>See local lab reference range</td>
<td>None</td>
</tr>
<tr>
<td>Mean corpuscular hemoglobin</td>
<td>See local lab reference range</td>
<td>None</td>
</tr>
<tr>
<td>Mean corpuscular hemoglobin content</td>
<td>See local lab reference range</td>
<td>None</td>
</tr>
</tbody>
</table>

*Lab calls Field Centers. Field center notifies participant and participant’s physician by telephone/fax if participant has granted permission to notify physician. Use modified letter from CHS (see Appendix 6) with abnormal value filled in.

**Notify participant and participant’s physician by fax/letter if participant has granted permission to notify physician. Use modified letter from CHS with abnormal value filled in.
8. Quality Assurance

8.1 Training Requirements

Clinical experience with phlebotomy is mandatory. Additional training should include:

- Read and study manual
- Attend Health ABC training session on techniques (or observe procedure by experienced examiner)
- Discuss problems and questions with local expert or QC officer

8.2 Certification Requirements

- Complete training requirements
- Explain what to do for difficult venipuncture
- Recite measures to take for fainting participant
- Conduct phlebotomy on volunteer or participant while being observed by QC officer using QC checklist

8.3 Quality Assurance Checklist

Preparation:
- Blood collection trays properly prepared
- Blood draw tubes properly labeled
- Questions on Phlebotomy form asked
- Hepatitis B vaccination given or offered to all personnel handling blood
Venipuncture properly carried out:
- Script properly delivered
- Non-permeable lab coats, gloves, and face shields used
- Preparation of venipuncture site correctly done
- Venipuncture smoothly done
- Tubes filled in proper priority order
- CPT tubes at least 1/2 full
- Tourniquet removed at 2 minutes
- Needle removed and arm bandaged correctly
- Needle and tubing appropriately disposed

Tubes mixed and handled correctly after filling:
- Tube 1 mixed gently, placed in ice bath
- Tubes 2 & 3 mixed gently, placed in rack at room temperature
- Tube 4 NOT mixed, placed in rack at room temperature

Phlebotomy form properly filled out:
- Sample ID barcode label affixed to upper right corner (and to lower left corner of Lab Processing form)
- Time at start of venipuncture entered (12-hr time, am/pm used)
- Time at end of venipuncture entered
- Total elapsed time with tourniquet entered
- Quality of venipuncture checked
- Question 9 filled out for all blood collection tubes
- Total fasting time correctly calculated
## APPENDIX 1 Sample Label Sheet (Bar Codes)
### (page 1 of 2)

| Draw Tube 1 | Draw Tube 2 | Draw Tube 3 | Backup
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Purple top 3 mL</td>
<td>Black/blue top 8 mL</td>
<td>Black/blue top 8 mL</td>
<td>Vacutainer</td>
</tr>
<tr>
<td>Draw Tube 4</td>
<td>Backup</td>
<td>Backup</td>
<td></td>
</tr>
<tr>
<td>Red top 10 mL</td>
<td>Vacutainer</td>
<td>Vacutainer</td>
<td></td>
</tr>
<tr>
<td>Backup</td>
<td>Phlebotomy Form</td>
<td>Laboratory Processing Form</td>
<td></td>
</tr>
<tr>
<td>BDID Form</td>
<td>BDID Form</td>
<td>BDID Form</td>
<td></td>
</tr>
</tbody>
</table>

### Placement Instructions:
- Place this end on vial first
- PD-01 R/Serum 1.5
- PD-02 R/Serum 1.5
- PD-03 R/Serum 1.5
- PD-04 R/Serum 1.5
- Citrate Plasma Pool 10 mL
- PD-05 B/Citrate 1.0 To LCBR
- PD-06 B/Citrate 1.0 To LCBR

### Additional Information:
- Pool 15 mL conical centrifuge tube
- Step 2: Cells
- Step 3: Platelets
- To LCBR
### APPENDIX 1
Sample Label Sheet (Bar Codes)
(page 2 of 2)

<table>
<thead>
<tr>
<th>Place this end on vial first</th>
<th>Place this end on vial first</th>
<th>Place this end on vial first</th>
</tr>
</thead>
<tbody>
<tr>
<td>#######-07 B/Citrate 1.0</td>
<td>#######-08 B/Citrate 1.0</td>
<td>#######-13 C/Buffy 2.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Use special freezing protocol</td>
</tr>
<tr>
<td>Place this end on vial first</td>
<td>Place this end on vial first</td>
<td>Place this end on vial first</td>
</tr>
<tr>
<td>#######-14 C/Buffy 2.0</td>
<td>#######-15 O/platelets 2.0</td>
<td>#######-16 O/platelets 2.0</td>
</tr>
<tr>
<td>Use special freezing protocol</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
APPENDIX 2 Phlebotomy Form

1. Did the participant have cells collected during Visit 2?  
(Examiner Note: Refer to Data from Prior Visits Report.)
   - Yes
   - No

   Do not fill CPT tubes.

2. Did participant refuse cell collection during Visit 2?  
(Examiner Note: Refer to Data from Prior Visit Report.)
   - Yes
   - No

   Do not fill CPT tubes.
   - Fill CPT tubes.
   - Do NOT combine draw with arterialized blood gas protocol.

3. Do you bleed or bruise easily?  
   - Yes
   - No
   - Don't know
   - Refused

4. Have you ever experienced fainting spells while having blood drawn?  
   - Yes
   - No
   - Don't know
   - Refused

5. Have you ever had a radical mastectomy? (Female Participants Only)  
   - Yes
   - No
   - Don't know
   - Refused

   Which side?  
   - Right
   - Left
   - Both

   Draw blood on left side.
   Draw blood on right side.
   Do not draw blood.

6. Have you ever had a graft for kidney dialysis?  
   - Yes
   - No
   - Don't know
   - Refused

   Which side?  
   - Right
   - Left
   - Both

   Draw blood on left side.
   Draw blood on right side.
   Do not draw blood.
PHLEBOTOMY

6. Is participant currently receiving supplemental oxygen?
   - Yes
   - No
   - Don't know
   - Refused

   How much?
   ____________________________ liters/min

7. Participant's temperature:
   ____________________________ ° F

8. Time at start of venipuncture?
   - Hours
   - Minutes
   - am
   - pm

   Please describe why not?
   ____________________________

9. Time blood draw completed:
   - Hours
   - Minutes
   - am
   - pm

10. Total tourniquet time:
    (Examiner Note: if tourniquet was reapplied, enter total time tourniquet was on. Note that 2 minutes is optimum.)
    ____________________________ minutes

   Comments on phlebotomy:
   ____________________________
   ____________________________

11. What is the date and time you last ate anything?
    a. Date of last food:
       / / Year
       Month     Day
    b. Time of last food:
       - Hours
       - Minutes
       - am
       - pm
    c. How many hours have passed since the participant last ate any food?
       ____________________________ hours (Question 9 minus Question 11b. Round to nearest hour)
## PHLEBOTOMY

### Quality of venipuncture:
- Clean
- Traumatic
- Vein collapse
- Hematoma
- Vein hard to get
- Multiple sticks
- Excessive duration of draw
- Leakage at venipuncture site
- Other (Please specify)

### Was arterialized venous blood sample obtained?
- Yes
- No

### Was a standard blood draw done?
- Yes
- No

### Were tubes filled to specified capacity? If not, comment why.

<table>
<thead>
<tr>
<th>Blood Volume/Tube</th>
<th>Filled to Capacity?</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. CBC 3 ml</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>2. CPT 8 ml</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>3. CPT 8 ml</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>4. Serum 10 ml</td>
<td>Yes</td>
<td></td>
</tr>
</tbody>
</table>

---

Page Link #   | *Page 44*
APPENDIX 3  Return Visit Laboratory Forms
(page 1 of 4)

Why did the participant return for phlebotomy?
(Examiner Note: Please mark all that apply.)

- For standard phlebotomy (cells and/or serum) only
- For arterialized venous blood sample only
- For both (either combined or separately)

Did the participant have cells collected during Visit 2?
(Examiner Note: Refer to Data from Prior Visits Report.)

- Yes
- No

Did participant refuse cell collection during Visit 2?
(Examiner Note: Refer to Data from Prior Visit Report.)

- Yes
- No

Do not fill CPT tubes.

- Fill CPT tubes.
- Do NOT combine draw with arterialized blood gas protocol.

Do you bleed or bruise easily?

- Yes
- No
- Don't know
- Refused

Have you ever experienced fainting spells while having blood drawn?

- Yes
- No
- Don't know
- Refused

Have you ever had a radical mastectomy? (Female Participants Only)

- Yes
- No
- Don't know
- Refused

Which side?

- Right
- Left
- Both

Draw blood on left side
Draw blood on right side
Do not draw blood.

LCBR Use only: Received Date: Time:

Page Link # Frozen? Page 1

Y3RVP Version 1.0, 7/9/99

Version 1.0
10/1/99
APPENDIX 3 Return Visit Laboratory Forms
(page 2 of 4)

**Health ABC YEAR 3 RETURN VISIT PHLEBOTOMY**

6. Have you ever had a graft for kidney dialysis?
   - Yes
   - No
   - Don't know
   - Refused

   Which side?
   - Right
   - Left
   - Both

   Draw blood on left side
   Draw blood on right side
   Do not draw blood.

7. Is participant currently receiving supplemental oxygen?
   - Yes
   - No
   - Don't know
   - Refused

   How much?
   liters/min

8. Participant's temperature:
   ° F

9. Time at start of venipuncture?
   Hours:
   Minutes:
   am
   pm

   a. Was any blood drawn?
      - Yes
      - No

   Please describe why not?

10. Time blood draw completed:
    Hours:
    Minutes:
    am
    pm

11. Total tourniquet time:
    (Examiner Note: If tourniquet was reapplied, enter total time tourniquet was on. Note that 2 minutes is optimum.)
    minutes

    Comments on phlebotomy:

---

Page Link #  •Page 2 •  Y3RVP Version 1.0, 7/9/99
APPENDIX 3 Return Visit Laboratory Forms (page 3 of 4)

**YEAR 3 RETURN VISIT PHLEBOTOMY**

12. What is the date and time you last ate anything?
   a. Date of last food: [ ] [ ] [ ]
      Month Day Year
   b. Time of last food: [ ] [: ]
      Hours Minutes
   c. How many hours have passed since the participant last ate any food? [ ] [ ] hours
      (Question 10 minus Question 12b. Round to nearest hour)

13. Quality of venipuncture:
   - Clean
   - Traumatic
     - Vein collapse
     - Excessive duration of draw
     - Hematoma
     - Leakage at venipuncture site
     - Vein hard to get
     - Other (Please specify:)
     - Multiple sticks

14. Was arterialized venous blood sample obtained?
   - Yes
   - No

15. Was a standard blood draw done?
   - Yes
   - No

16. Were tubes filled to specified capacity? If not, comment why.

<table>
<thead>
<tr>
<th>Blood Volume/Tube</th>
<th>Filled to Capacity?</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. CBC 3 ml</td>
<td>[ ] Yes [ ] No</td>
<td></td>
</tr>
<tr>
<td>2. CPT 8 ml</td>
<td>[ ] Yes [ ] No</td>
<td></td>
</tr>
<tr>
<td>3. CPT 8 ml</td>
<td>[ ] Yes [ ] No</td>
<td></td>
</tr>
<tr>
<td>4. Serum 10 ml</td>
<td>[ ] Yes [ ] No</td>
<td></td>
</tr>
</tbody>
</table>
### APPENDIX 3  Return Visit Laboratory Forms
(page 4 of 4)

**YEAR 3 RETURN VISIT BLOOD GAS**

1. Controls (yellow)
   - pH
   - pCO₂
   - pO₂

2. Participant's blood gas:
   - pH
   - pCO₂
   - pO₂
   - HCO₃

3. Controls (red)
   - pH
   - pCO₂
   - pO₂

4. Were controls in range?
   - Yes
   - No

5. Room temperature where visits are stored:
   - °C

**YEAR 3 RETURN VISIT LABORATORY**

<table>
<thead>
<tr>
<th>Collection Tubes</th>
<th>Cryo Vol. Type</th>
<th>To Fill in bubble</th>
<th>Problems</th>
<th>Staff ID #</th>
</tr>
</thead>
<tbody>
<tr>
<td>#4 Serum</td>
<td>01 1.0 R/1.5 M</td>
<td>O</td>
<td>H</td>
<td>P</td>
</tr>
<tr>
<td></td>
<td>02 1.0 R/1.5 M</td>
<td>O</td>
<td>H</td>
<td>P</td>
</tr>
<tr>
<td></td>
<td>03 1.0 R/1.5 M</td>
<td>O</td>
<td>H</td>
<td>P</td>
</tr>
<tr>
<td></td>
<td>04 1.0 R/1.5 M</td>
<td>O</td>
<td>H</td>
<td>P</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Collection Tubes</th>
<th>Cryo Vol. Type</th>
<th>To Fill in bubble</th>
<th>Problems</th>
<th>Staff ID #</th>
</tr>
</thead>
<tbody>
<tr>
<td>#2, 3 Citrate</td>
<td>05 1.0 B/1.5 L</td>
<td>O</td>
<td>H</td>
<td>P</td>
</tr>
<tr>
<td></td>
<td>06 1.0 B/1.5 L</td>
<td>O</td>
<td>H</td>
<td>P</td>
</tr>
<tr>
<td></td>
<td>07 1.0 B/1.5 L</td>
<td>O</td>
<td>H</td>
<td>P</td>
</tr>
<tr>
<td></td>
<td>08 1.0 B/1.5 L</td>
<td>O</td>
<td>H</td>
<td>P</td>
</tr>
</tbody>
</table>

- M=McKesson; H=Hemolyzed; P=Partia; R=Red; C=Clear; B=Blue; O=Orange
- *Place in a styrofoam box at -20°C for 2 hours. Transfer to -80°C to hold for shipping.*
APPENDIX 4 OSHA Guidelines

Enforcement Procedures for the Occupational Exposure to Bloodborne Pathogens
Standard, 29 CFR 1910.1030
(15 pages)

1910.1030 - Bloodborne pathogens.

* Standard Number: 1910.1030
* Standard Title: Bloodborne pathogens.
* SubPart Number: Z
* SubPart Title: Toxic and Hazardous Substances

Produced by USDOL OSHA - Directorate of Safety Standards &
Directorate of Health Standards
Maintained by USDOL OSHA - OCIS

(a) Scope and Application. This section applies to all occupational exposure to blood or other potentially infectious materials as defined by paragraph (b) of this section.

(b) Definitions. For purposes of this section, the following shall apply:

"Assistant Secretary" means the Assistant Secretary of Labor for Occupational Safety and Health, or designated representative.
"Blood" means human blood, human blood components, and products made from human blood.
"Bloodborne Pathogens" means pathogenic microorganisms that are present in human blood and can cause disease in humans. These pathogens include, but are not limited to, hepatitis B virus (HBV) and human immunodeficiency virus (HIV).
"Clinical Laboratory" means a workplace where diagnostic or other screening procedures are performed on blood or other potentially infectious materials.
"Contaminated" means the presence or the reasonably anticipated presence of blood or other potentially infectious materials on an item or surface.
"Contaminated Laundry" means laundry which has been soiled with blood or
other potentially infectious materials or may contain sharps.

"Contaminated Sharps" means any contaminated object that can penetrate the skin including, but not limited to, needles, scalpels, broken glass, broken capillary tubes, and exposed ends of dental wires.

"Decontamination" means the use of physical or chemical means to remove, inactivate, or destroy bloodborne pathogens on a surface or item to the point where they are no longer capable of transmitting infectious particles and the surface or item is rendered safe for handling, use, or disposal.

"Director" means the Director of the National Institute for Occupational Safety and Health, U.S. Department of Health and Human Services, or designated representative.

"Engineering Controls" means controls (e.g., sharps disposal containers, self-sheathing needles) that isolate or remove the bloodborne pathogens hazard from the workplace.

"Exposure Incident" means a specific eye, mouth, other mucous membrane, non-intact skin, or parenteral contact with blood or other potentially infectious materials that results from the performance of an employee's duties.

"Handwashing Facilities" means a facility providing an adequate supply of running potable water, soap and single use towels or hot air drying machines.

"Licensed Healthcare Professional" is a person whose legally permitted scope of practice allows him or her to independently perform the activities required by paragraph (f) Hepatitis B Vaccination and Post-exposure Evaluation and Follow-up.

"HBV" means hepatitis B virus.

"HIV" means human immunodeficiency virus.

"Occupational Exposure" means reasonably anticipated skin, eye, mucous membrane, or parenteral contact with blood or other potentially infectious materials that may result from the performance of an employee's duties.

"Other Potentially Infectious Materials" means (1) The following human body fluids: semen, vaginal secretions, cerebrospinal fluid, synovial fluid, pleural fluid, pericardial fluid, peritoneal fluid, amniotic fluid, saliva in dental procedures, any body fluid that is visibly contaminated with blood, and all body fluids in situations where it is difficult or impossible to differentiate between body fluids; (2) Any unfixed tissue or organ (other than intact skin) from a human (living or dead); and (3) HIV-containing cell or tissue cultures, organ cultures, and HIV- or HBV-containing culture medium or other solutions; and blood, organs, or
other tissues from experimental animals infected with HIV or HBV.

"Parenteral" means piercing mucous membranes or the skin barrier through such events as needlesticks, human bites, cuts, and abrasions.

"Personal Protective Equipment" is specialized clothing or equipment worn by an employee for protection against a hazard. General work clothes (e.g., uniforms, pants, shirts or blouses) not intended to function as protection against a hazard are not considered to be personal protective equipment.

"Production Facility" means a facility engaged in industrial-scale, large-volume or high concentration production of HIV or HBV.

"Regulated Waste" means liquid or semi-liquid blood or other potentially infectious materials; contaminated items that would release blood or other potentially infectious materials in a liquid or semi-liquid state if compressed; items that are caked with dried blood or other potentially infectious materials and are capable of releasing these materials during handling; contaminated sharps; and pathological and microbiological wastes containing blood or other potentially infectious materials.

"Research Laboratory" means a laboratory producing or using research-laboratory-scale amounts of HIV or HBV. Research laboratories may produce high concentrations of HIV or HBV but not in the volume found in production facilities.

"Source Individual" means any individual, living or dead, whose blood or other potentially infectious materials may be a source of occupational exposure to the employee. Examples include, but are not limited to, hospital and clinic patients; clients in institutions for the developmentally disabled; trauma victims; clients of drug and alcohol treatment facilities; residents of hospices and nursing homes; human remains; and individuals who donate or sell blood or blood components.

"Sterilize" means the use of a physical or chemical procedure to destroy all microbial life including highly resistant bacterial endospores.

"Universal Precautions" is an approach to infection control. According to the concept of Universal Precautions, all human blood and certain human body fluids are treated as if known to be infectious for HIV, HBV, and other bloodborne pathogens.

"Work Practice Controls" means controls that reduce the likelihood of exposure by altering the manner in which a task is performed (e.g., prohibiting recapping of needles by a two-handed technique).

..1910.1030(c) (c) Exposure Control. (1) Exposure Control Plan. (i) Each employer having an employee(s) with occupational exposure as defined by
paragraph (b) of this section shall establish a written Exposure Control Plan designed to eliminate or minimize employee exposure. (ii) The Exposure Control Plan shall contain at least the following elements: (A) The exposure determination required by paragraph (c)(2), (B) The schedule and method of implementation for paragraphs (d) Methods of Compliance, (e) HIV and HBV Research Laboratories and Production Facilities, (f) Hepatitis B Vaccination and Post-Exposure Evaluation and Follow-up, (g) Communication of Hazards to Employees, and (h) Recordkeeping, of this standard, and (C) The procedure for the evaluation of circumstances surrounding exposure incidents as required by paragraph (f)(3)(i) of this standard. (iii) Each employer shall ensure that a copy of the Exposure Control Plan is accessible to employees in accordance with 29 CFR 1910.1020(e). (iv) The Exposure Control Plan shall be reviewed and updated at least annually and whenever necessary to reflect new or modified tasks and procedures which affect occupational exposure and to reflect new or revised employee positions with occupational exposure.  

.1910.1030(c)(1)(v) (v) The Exposure Control Plan shall be made available to the Assistant Secretary and the Director upon request for examination and copying. (2) Exposure Determination. (i) Each employer who has an employee(s) with occupational exposure as defined by paragraph (b) of this section shall prepare an exposure determination. This exposure determination shall contain the following: (A) A list of all job classifications in which all employees in those job classifications have occupational exposure; (B) A list of job classifications in which some employees have occupational exposure, and (C) A list of all tasks and procedures or groups of closely related task and procedures in which occupational exposure occurs and that are performed by employees in job classifications listed in accordance with the provisions of paragraph (c)(2)(i)(B) of this standard. (ii) This exposure determination shall be made without regard to the use of personal protective equipment. (d) Methods of Compliance. (1) General. Universal precautions shall be observed to prevent contact with blood or other potentially infectious materials. Under circumstances in which differentiation between body fluid types is difficult or impossible, all body fluids shall be considered potentially infectious materials.

.1910.1030(d)(2) (2) Engineering and Work Practice Controls. (i) Engineering and work practice controls shall be used to eliminate or minimize employee exposure. Where occupational exposure remains after institution of these controls, personal protective equipment shall also be used. (ii) Engineering controls shall be examined and maintained or replaced on a regular schedule to ensure their effectiveness. (iii) Employers shall provide handwashing facilities which are readily accessible to employees.
(iv) When provision of handwashing facilities is not feasible, the employer shall provide either an appropriate antiseptic hand cleanser in conjunction with clean cloth/ paper towels or antiseptic towelettes. When antiseptic hand cleansers or towelettes are used, hands shall be washed with soap and running water as soon as feasible. (v) Employers shall ensure that employees wash their hands immediately or as soon as feasible after removal of gloves or other personal protective equipment. (vi) Employers shall ensure that employees wash hands and any other skin with soap and water, or flush mucous membranes with water immediately or as soon as feasible following contact of such body areas with blood or other potentially infectious materials. 

..1910.1030(d)(2)(vii) (vii) Contaminated needles and other contaminated sharps shall not be bent, recapped, or removed except as noted in paragraphs (d)(2)(vii)(A) and (d)(2)(vii)(B) below. Shearing or breaking of contaminated needles is prohibited. (A) Contaminated needles and other contaminated sharps shall not be bent, recapped or removed unless the employer can demonstrate that no alternative is feasible or that such action is required by a specific medical or dental procedure. (B) Such bending, recapping or needle removal must be accomplished through the use of a mechanical device or a one-handed technique. (viii) Immediately or as soon as possible after use, contaminated reusable sharps shall be placed in appropriate containers until properly reprocessed. These containers shall be:

(A) puncture resistant;

(B) labeled or color-coded in accordance with this standard;

..1910.1030(d)(2)(viii)(C)

(C) leakproof on the sides and bottom; and

(D) in accordance with the requirements set forth in paragraph (d)(4)(ii)(E) for reusable sharps.

(ix) Eating, drinking, smoking, applying cosmetics or lip balm, and handling contact lenses are prohibited in work areas where there is a reasonable likelihood of occupational exposure. (x) Food and drink shall not be kept in refrigerators, freezers, shelves, cabinets or on countertops or benchtops where blood or other potentially infectious materials are present. (xi) All procedures involving blood or other potentially infectious materials shall be performed in such a manner as to minimize splashing, spraying,
spattering, and generation of droplets of these substances. (xii) Mouth pipetting/suctioning of blood or other potentially infectious materials is prohibited. (xiii) Specimens of blood or other potentially infectious materials shall be placed in a container which prevents leakage during collection, handling, processing, storage, transport, or shipping.

1910.1030(d)(2)(xiii)(A) (A) The container for storage, transport, or shipping shall be labeled or color-coded according to paragraph (g)(1)(i) and closed prior to being stored, transported, or shipped. When a facility utilizes Universal Precautions in the handling of all specimens, the labeling/color-coding of specimens is not necessary provided containers are recognizable as containing specimens. This exemption only applies while such specimens/containers remain within the facility. Labeling or color-coding in accordance with paragraph (g)(1)(i) is required when such specimens/containers leave the facility. (B) If outside contamination of the primary container occurs, the primary container shall be placed within a second container which prevents leakage during handling, processing, storage, transport, or shipping and is labeled or color-coded according to the requirements of this standard. (C) If the specimen could puncture the primary container, the primary container shall be placed within a secondary container which is puncture-resistant in addition to the above characteristics. (xiv) Equipment which may become contaminated with blood or other potentially infectious materials shall be examined prior to servicing or shipping and shall be decontaminated as necessary, unless the employer can demonstrate that decontamination of such equipment or portions of such equipment is not feasible. 1910.1030(d)(2)(xiv)(A) (A) A readily observable label in accordance with paragraph (g)(1)(i)(H) shall be attached to the equipment stating which portions remain contaminated. (B) The employer shall ensure that this information is conveyed to all affected employees, the servicing representative, and/or the manufacturer, as appropriate, prior to handling, servicing, or shipping so that appropriate precautions will be taken.

(3) Personal Protective Equipment. (i) Provision. When there is occupational exposure, the employer shall provide, at no cost to the employee, appropriate personal protective equipment such as, but not limited to, gloves, gowns, laboratory coats, face shields or masks and eye protection, and mouthpieces, resuscitation bags, pocket masks, or other ventilation devices. Personal protective equipment will be considered "appropriate" only if it does not permit blood or other potentially infectious materials to pass through to or reach the employee's work clothes, street clothes, undergarments, skin, eyes, mouth, or other mucous membranes under normal conditions of use and for the duration of time which
the protective equipment will be used. 1910.1030(d)(3)(ii) (ii) Use. The employer shall ensure that the employee uses appropriate personal protective equipment unless the employer shows that the employee temporarily and briefly declined to use personal protective equipment when, under rare and extraordinary circumstances, it was the employee's professional judgment that in the specific instance its use would have prevented the delivery of health care or public safety services or would have posed an increased hazard to the safety of the worker or co-worker. When the employee makes this judgment, the circumstances shall be investigated and documented in order to determine whether changes can be instituted to prevent such occurrences in the future. (iii) Accessibility. The employer shall ensure that appropriate personal protective equipment in the appropriate sizes is readily accessible at the worksite or is issued to employees. Hypoallergenic gloves, glove liners, powderless gloves, or other similar alternatives shall be readily accessible to those employees who are allergic to the gloves normally provided. (iv) Cleaning, Laundering, and Disposal. The employer shall clean, launder, and dispose of personal protective equipment required by paragraphs (d) and (e) of this standard, at no cost to the employee. 1910.1030(d)(3)(v) (v) Repair and Replacement. The employer shall repair or replace personal protective equipment as needed to maintain its effectiveness, at no cost to the employee. (vi) If a garment(s) is penetrated by blood or other potentially infectious materials, the garment(s) shall be removed immediately or as soon as feasible. (vii) All personal protective equipment shall be removed prior to leaving the work area. (viii) When personal protective equipment is removed it shall be placed in an appropriately designated area or container for storage, washing, decontamination or disposal. (ix) Gloves. Gloves shall be worn when it can be reasonably anticipated that the employee may have hand contact with blood, other potentially infectious materials, mucus membranes, and non-intact skin; when performing vascular access procedures except as specified in paragraph (d)(3)(ix)(D); and when handling or touching contaminated items or surfaces. (A) Disposable (single use) gloves such as surgical or examination gloves, shall be replaced as soon as practical when contaminated or as soon as feasible if they are torn, punctured, or when their ability to function as a barrier is compromised. 1910.1030(d)(3)(ix)(B) (B) Disposable (single use) gloves shall not be washed or decontaminated for re-use. (C) Utility gloves may be decontaminated for re-use if the integrity of the glove is not compromised. However, they must be discarded if they are cracked, peeling, torn, punctured, or exhibit other signs of deterioration or when their ability to
function as a barrier is compromised. (D) If an employer in a volunteer blood donation center judges that routine gloving for all phlebotomies is not necessary then the employer shall: {1} Periodically reevaluate this policy; {2} Make gloves available to all employees who wish to use them for phlebotomy; {3} Not discourage the use of gloves for phlebotomy; and {4} Require that gloves be used for phlebotomy in the following circumstances: [i] When the employee has cuts, scratches, or other breaks in his or her skin; [ii] When the employee judges that hand contamination with blood may occur, for example, when performing phlebotomy on an uncooperative source individual; and [iii] When the employee is receiving training in phlebotomy. ..1910.1030(d)(3)(x) (x) Masks, Eye Protection, and Face Shields. Masks in combination with eye protection devices, such as goggles or glasses with solid side shields, or chin-length face shields, shall be worn whenever splashes, spray, spatter, or droplets of blood or other potentially infectious materials may be generated and eye, nose, or mouth contamination can be reasonably anticipated. (xi) Gowns, Aprons, and Other Protective Body Clothing. Appropriate protective clothing such as, but not limited to, gowns, aprons, lab coats, clinic jackets, or similar outer garments shall be worn in occupational exposure situations. The type and characteristics will depend upon the task and degree of exposure anticipated. (xii) Surgical caps or hoods and/ or shoe covers or boots shall be worn in instances when gross contamination can reasonably be anticipated (e.g., autopsies, orthopaedic surgery). (4) Housekeeping. (i) General. Employers shall ensure that the worksite is maintained in a clean and sanitary condition. The employer shall determine and implement an appropriate written schedule for cleaning and method of decontamination based upon the location within the facility, type of surface to be cleaned, type of soil present, and tasks or procedures being performed in the area. ..1910.1030(d)(4)(ii) (ii) All equipment and environmental and working surfaces shall be cleaned and decontaminated after contact with blood or other potentially infectious materials. (A) Contaminated work surfaces shall be decontaminated with an appropriate disinfectant after completion of procedures; immediately or as soon as feasible when surfaces are overtly contaminated or after any spill of blood or other potentially infectious materials; and at the end of the work shift if the surface may have become contaminated since the last cleaning. (B) Protective coverings, such as plastic wrap, aluminum foil, or imperviously-backed absorbent paper used to cover equipment and environmental surfaces, shall be removed and replaced as soon as feasible when they become overtly contaminated or at the end of the workshift if they may have become contaminated during the shift. (C) All bins, pails, cans,
and similar receptacles intended for reuse which have a reasonable likelihood for becoming contaminated with blood or other potentially infectious materials shall be inspected and decontaminated on a regularly scheduled basis and cleaned and decontaminated immediately or as soon as feasible upon visible contamination. 

1910.1030(d)(4)(ii)(D) (D) Broken glassware which may be contaminated shall not be picked up directly with the hands. It shall be cleaned up using mechanical means, such as a brush and dust pan, tongs, or forceps. (E) Reusable sharps that are contaminated with blood or other potentially infectious materials shall not be stored or processed in a manner that requires employees to reach by hand into the containers where these sharps have been placed. (iii) Regulated Waste. (A) Contaminated Sharps Discarding and Containment. (1) Contaminated sharps shall be discarded immediately or as soon as feasible in containers that are: [a] Closable; [b] Puncture resistant; [c] Leakproof on sides and bottom; and [d] Labeled or color-coded in accordance with paragraph (g)(1)(i) of this standard. (2) During use, containers for contaminated sharps shall be: [a] Easily accessible to personnel and located as close as is feasible to the immediate area where sharps are used or can be reasonably anticipated to be found (e.g., laundries); [b] Maintained upright throughout use; and [c] Replaced routinely and not be allowed to overfill.

1910.1030(d)(4)(iii)(A) When moving containers of contaminated sharps from the area of use, the containers shall be: [a] Closed immediately prior to removal or replacement to prevent spillage or protrusion of contents during handling, storage, transport, or shipping; [b] Placed in a secondary container if leakage is possible. The second container shall be: [i] Closable; [ii] Constructed to contain all contents and prevent leakage during handling, storage, transport, or shipping; and [iii] Labeled or color-coded according to paragraph (g)(1)(i) of this standard. (4) Reusable containers shall not be opened, emptied, or cleaned manually or in any other manner which would expose employees to the risk of percutaneous injury. (B) Other Regulated Waste Containment. (1) Regulated waste shall be placed in containers which are: [a] Closable; [b] Constructed to contain all contents and prevent leakage of fluids during handling, storage, transport or shipping; [c] Labeled or color-coded in accordance with paragraph (g)(1)(i) this standard; and [d] Closed prior to removal to prevent spillage or protrusion of contents during handling, storage, transport, or shipping.

1910.1030(d)(4)(iii)(B) If outside contamination of the regulated waste container occurs, it shall be placed in a second container. The second container shall be: [a] Closable; [b] Constructed to contain all contents and prevent leakage of fluids during handling, storage, transport or
shipping; [c] Labeled or color-coded in accordance with paragraph (g)(1)(i) of this standard; and [d] Closed prior to removal to prevent spillage or protrusion of contents during handling, storage, transport, or shipping. (C) Disposal of all regulated waste shall be in accordance with applicable regulations of the United States, States and Territories, and political subdivisions of States and Territories. (iv) Laundry. (A) Contaminated laundry shall be handled as little as possible with a minimum of agitation. (1) Contaminated laundry shall be bagged or containerized at the location where it was used and shall not be sorted or rinsed in the location of use. *1910.1030(d)(4)(iv)(A) {2} *2 Contaminated laundry shall be placed and transported in bags or containers labeled or color-coded in accordance with paragraph (g)(1)(i) of this standard. When a facility utilizes Universal Precautions in the handling of all soiled laundry, alternative labeling or color-coding is sufficient if it permits all employees to recognize the containers as requiring compliance with Universal Precautions. (2) Whenever contaminated laundry is wet and presents a reasonable likelihood of soak-through of or leakage from the bag or container, the laundry shall be placed and transported in bags or containers which prevent soak-through and/ or leakage of fluids to the exterior. (B) The employer shall ensure that employees who have contact with contaminated laundry wear protective gloves and other appropriate personal protective equipment. (C) When a facility ships contaminated laundry off-site to a second facility which does not utilize Universal Precautions in the handling of all laundry, the facility generating the contaminated laundry must place such laundry in bags or containers which are labeled or color-coded in accordance with paragraph (g)(1)(i). *1910.1030(e) (e) HIV and HBV Research Laboratories and Production Facilities. (1) This paragraph applies to research laboratories and production facilities engaged in the culture, production, concentration, experimentation, and manipulation of HIV and HBV. It does not apply to clinical or diagnostic laboratories engaged solely in the analysis of blood, tissues, or organs. These requirements apply in addition to the other requirements of the standard. (2) Research laboratories and production facilities shall meet the following criteria: (i) Standard Microbiological Practices. All regulated waste shall either be incinerated or decontaminated by a method such as autoclaving known to effectively destroy bloodborne pathogens. (ii) Special Practices (A) Laboratory doors shall be kept closed when work involving HIV or HBV is in progress. (B) Contaminated materials that are to be decontaminated at a site away from the work area shall be placed in a durable, leakproof, labeled or color-coded container that is closed before being removed from the work area. *1910.1030(e)(2)(ii)(C)
Access to the work area shall be limited to authorized persons. Written policies and procedures shall be established whereby only persons who have been advised of the potential biohazard, who meet any specific entry requirements, and who comply with all entry and exit procedures shall be allowed to enter the work areas and animal rooms. (D) When other potentially infectious materials or infected animals are present in the work area or containment module, a hazard warning sign incorporating the universal biohazard symbol shall be posted on all access doors. The hazard warning sign shall comply with paragraph (g)(1)(ii) of this standard. (E) All activities involving other potentially infectious materials shall be conducted in biological safety cabinets or other physical-containment devices within the containment module. No work with these other potentially infectious materials shall be conducted on the open bench. (F) Laboratory coats, gowns, smocks, uniforms, or other appropriate protective clothing shall be used in the work area and animal rooms. Protective clothing shall not be worn outside of the work area and shall be decontaminated before being laundered. (G) Special care shall be taken to avoid skin contact with other potentially infectious materials. Gloves shall be worn when handling infected animals and when making hand contact with other potentially infectious materials is unavoidable. (H) Before disposal all waste from work areas and from animal rooms shall either be incinerated or decontaminated by a method such as autoclaving known to effectively destroy bloodborne pathogens. (I) Vacuum lines shall be protected with liquid disinfectant traps and high-efficiency particulate air (HEPA) filters or filters of equivalent or superior efficiency and which are checked routinely and maintained or replaced as necessary. (J) Hypodermic needles and syringes shall be used only for parenteral injection and aspiration of fluids from laboratory animals and diaphragm bottles. Only needle-locking syringes or disposable syringe-needle units (i.e., the needle is integral to the syringe) shall be used for the injection or aspiration of other potentially infectious materials. Extreme caution shall be used when handling needles and syringes. A needle shall not be bent, sheared, replaced in the sheath or guard, or removed from the syringe following use. The needle and syringe shall be promptly placed in a puncture-resistant container and autoclaved or decontaminated before reuse or disposal. (K) All spills shall be immediately contained and cleaned up by appropriate professional staff or others properly trained and equipped to work with potentially concentrated infectious materials. (L) A spill or accident that results in an exposure incident shall be immediately reported to the laboratory director or other responsible person.
biosafety manual shall be prepared or adopted and periodically reviewed and updated at least annually or more often if necessary. Personnel shall be advised of potential hazards, shall be required to read instructions on practices and procedures, and shall be required to follow them. (iii) Containment Equipment. (A) Certified biological safety cabinets (Class I, II, or III) or other appropriate combinations of personal protection or physical containment devices, such as special protective clothing, respirators, centrifuge safety cups, sealed centrifuge rotors, and containment caging for animals, shall be used for all activities with other potentially infectious materials that pose a threat of exposure to droplets, splashes, spills, or aerosols. ..1910.1030(e)(2)(iii)(B) (B) Biological safety cabinets shall be certified when installed, whenever they are moved and at least annually. (3) HIV and HBV research laboratories shall meet the following criteria: (i) Each laboratory shall contain a facility for hand washing and an eye wash facility which is readily available within the work area. (ii) An autoclave for decontamination of regulated waste shall be available. (4) HIV and HBV production facilities shall meet the following criteria: (i) The work areas shall be separated from areas that are open to unrestricted traffic flow within the building. Passage through two sets of doors shall be the basic requirement for entry into the work area from access corridors or other contiguous areas. Physical separation of the high-containment work area from access corridors or other areas or activities may also be provided by a double-doored clothes-change room (showers may be included), airlock, or other access facility that requires passing through two sets of doors before entering the work area. ..1910.1030(e)(4)(ii) (ii) The surfaces of doors, walls, floors and ceilings in the work area shall be water resistant so that they can be easily cleaned. Penetrations in these surfaces shall be sealed or capable of being sealed to facilitate decontamination. (iii) Each work area shall contain a sink for washing hands and a readily available eye wash facility. The sink shall be foot, elbow, or automatically operated and shall be located near the exit door of the work area. (iv) Access doors to the work area or containment module shall be self-closing. (v) An autoclave for decontamination of regulated waste shall be available within or as near as possible to the work area. (vi) A ducted exhaust-air ventilation system shall be provided. This system shall create directional airflow that draws air into the work area through the entry area. The exhaust air shall not be recirculated to any other area of the building, shall be discharged to the outside, and shall be dispersed away from occupied areas and air intakes. The proper direction of the airflow shall be verified (i.e., into the work area...
area). ..1910.1030(e)(5) (5) Training Requirements. Additional training requirements for employees in HIV and HBV research laboratories and HIV and HBV production facilities are specified in paragraph (g)(2)(ix). (f) Hepatitis B Vaccination and Post-exposure Evaluation and Follow-up. (1) General. (i) The employer shall make available the hepatitis B vaccine and vaccination series to all employees who have occupational exposure, and post-exposure evaluation and follow-up to all employees who have had an exposure incident. (ii) The employer shall ensure that all medical evaluations and procedures including the hepatitis B vaccine and vaccination series and post-exposure evaluation and follow-up, including prophylaxis, are: (A) Made available at no cost to the employee; (B) Made available to the employee at a reasonable time and place; (C) Performed by or under the supervision of a licensed physician or by or under the supervision of another licensed healthcare professional; and (D) Provided according to recommendations of the U.S. Public Health Service current at the time these evaluations and procedures take place, except as specified by this paragraph (f). ..1910.1030(f)(1)(iii) (iii) The employer shall ensure that all laboratory tests are conducted by an accredited laboratory at no cost to the employee. (2) Hepatitis B Vaccination. (i) Hepatitis B vaccination shall be made available after the employee has received the training required in paragraph (g)(2)(vii)(I) and within 10 working days of initial assignment to all employees who have occupational exposure unless the employee has previously received the complete hepatitis B vaccination series, antibody testing has revealed that the employee is immune, or the vaccine is contraindicated for medical reasons. (ii) The employer shall not make participation in a prescreening program a prerequisite for receiving hepatitis B vaccination. (iii) If the employee initially declines hepatitis B vaccination but at a later date while still covered under the standard decides to accept the vaccination, the employer shall make available hepatitis B vaccination at that time. (iv) The employer shall assure that employees who decline to accept hepatitis B vaccination offered by the employer sign the statement in Appendix A. ..1910.1030(f)(2)(v) (v) If a routine booster dose(s) of hepatitis B vaccine is recommended by the U.S. Public Health Service at a future date, such booster dose(s) shall be made available in accordance with section (f)(1)(ii). (3) Post-exposure Evaluation and Follow-up. Following a report of an exposure incident, the employer shall make immediately available to the exposed employee a confidential medical evaluation and follow-up, including at least the following elements: (i) Documentation of the route(s) of exposure, and the circumstances under which the exposure incident occurred; (ii)
Identification and documentation of the source individual, unless the employer can establish that identification is infeasible or prohibited by state or local law; (A) The source individual’s blood shall be tested as soon as feasible and after consent is obtained in order to determine HBV and HIV infectivity. If consent is not obtained, the employer shall establish that legally required consent cannot be obtained. When the source individual’s consent is not required by law, the source individual’s blood, if available, shall be tested and the results documented.

(B) When the source individual is already known to be infected with HBV or HIV, testing for the source individual’s known HBV or HIV status need not be repeated. (C) Results of the source individual’s testing shall be made available to the exposed employee, and the employee shall be informed of applicable laws and regulations concerning disclosure of the identity and infectious status of the source individual. (iii)

Collection and testing of blood for HBV and HIV serological status; (A) The exposed employee’s blood shall be collected as soon as feasible and tested after consent is obtained. (B) If the employee consents to baseline blood collection, but does not give consent at that time for HIV serologic testing, the sample shall be preserved for at least 90 days. If, within 90 days of the exposure incident, the employee elects to have the baseline sample tested, such testing shall be done as soon as feasible. (iv)

Post-exposure prophylaxis, when medically indicated, as recommended by the U.S. Public Health Service; (v) Counseling; and (vi)

Evaluation of reported illnesses. (4) Information Provided to the Healthcare Professional. (i) The employer shall ensure that the healthcare professional responsible for the employee’s Hepatitis B vaccination is provided a copy of this regulation. (ii) The employer shall ensure that the healthcare professional evaluating an employee after an exposure incident is provided the following information: (A) A copy of this regulation; (B) A description of the exposed employee’s duties as they relate to the exposure incident; (C) Documentation of the route(s) of exposure and circumstances under which exposure occurred; (D) Results of the source individual’s blood testing, if available; and (E) All medical records relevant to the appropriate treatment of the employee including vaccination status which are the employer’s responsibility to maintain. (5) Healthcare Professional’s Written Opinion. The employer shall obtain and provide the employee with a copy of the evaluating healthcare professional’s written opinion within 15 days of the completion of the evaluation. (i) The healthcare professional’s written opinion for Hepatitis B vaccination shall be limited to whether Hepatitis B vaccination is indicated for an employee, and if the
employee has received such vaccination. (ii) The healthcare professional's written opinion for post-exposure evaluation and follow-up shall be limited to the following information: (A) That the employee has been informed of the results of the evaluation; and (B) That the employee has been told about any medical conditions resulting from exposure to blood or other potentially infectious materials which require further evaluation or treatment. (iii) All other findings or diagnoses shall remain confidential and shall not be included in the written report. (6) Medical Recordkeeping. Medical records required by this standard shall be maintained in accordance with paragraph (h)(1) of this section. (g) Communication of Hazards to Employees. (1) Labels and Signs. (i) Labels. (A) Warning labels shall be affixed to containers of regulated waste, refrigerators and freezers containing blood or other potentially infectious material; and other containers used to store, transport or ship blood or other potentially infectious materials, except as provided in paragraph (g)(1)(i)(E), (F) and (G).

..1910.1030(g)(1)(i)(B) (B) Labels required by this section shall include the following legend:

BIOHAZARD

(C) These labels shall be fluorescent orange or orange-red or predominantly so, with lettering and symbols in a contrasting color. (D) Labels shall be affixed as close as feasible to the container by string, wire, adhesive, or other method that prevents their loss or unintentional removal. (E) Red bags or red containers may be substituted for labels. (F) Containers of blood, blood components, or blood products that are labeled as to their contents and have been released for transfusion or other clinical use are exempted from the labeling requirements of paragraph (g). ..1910.1030(g)(1)(i)(G) (G) Individual containers of blood or other potentially infectious materials that are placed in a labeled container during storage, transport, shipment or disposal are exempted from the labeling requirement. (H) Labels required for contaminated equipment shall be in accordance with this paragraph and shall also state which portions of the equipment remain contaminated. (I) Regulated waste that has been decontaminated need not be labeled or color-coded. (ii) Signs. (A) The employer shall post signs at the entrance to work areas specified in paragraph (e), HIV and HBV Research Laboratory and Production Facilities, which shall bear the following legend:

BIOHAZARD
(Name of the Infectious Agent)
(Special requirements for entering the area)
(Name, telephone number of the laboratory director or other responsible person.)

..1910.1030(g)(1)(ii)(B) (B) These signs shall be fluorescent orange-red or predominantly so, with lettering and symbols in a contrasting color. (2) Information and Training. (i) Employers shall ensure that all employees with occupational exposure participate in a training program which must be provided at no cost to the employee and during working hours. (ii) Training shall be provided as follows: (A) At the time of initial assignment to tasks where occupational exposure may take place; (B) Within 90 days after the effective date of the standard; and (C) At least annually thereafter. (iii) For employees who have received training on bloodborne pathogens in the year preceding the effective date of the standard, only training with respect to the provisions of the standard which were not included need be provided. (iv) Annual training for all employees shall be provided within one year of their previous training. (v) Employers shall provide additional training when changes such as modification of tasks or procedures or institution of new tasks or procedures affect the employee's occupational exposure. The additional training may be limited to addressing the new exposures created. (vi) Material appropriate in content and vocabulary to educational level, literacy, and language of employees shall be used. (vii) The training program shall contain at least the following elements: (A) An accessible copy of the regulatory text of this standard and an explanation of its contents; (B) A general explanation of the epidemiology and symptoms of bloodborne diseases; (C) An explanation of the modes of transmission of bloodborne pathogens; (D) An explanation of the employer's exposure control plan and the means by which the employee can obtain a copy of the written plan; (E) An explanation of the appropriate methods for recognizing tasks and other activities that may involve exposure to blood and other potentially infectious materials; (F) An explanation of the use and limitations of methods that will prevent or reduce exposure including appropriate engineering controls, work practices, and personal protective equipment; (G) Information on the types, proper use, location, removal, handling, decontamination and disposal of personal protective equipment; (H) An explanation of the basis for selection of personal protective equipment; (I) Information on the hepatitis B vaccine, including information on its efficacy, safety, method of administration, the benefits of being vaccinated, and that the vaccine and vaccination will be
offered free of charge; (J) Information on the appropriate actions to take and persons to contact in an emergency involving blood or other potentially infectious materials; (K) An explanation of the procedure to follow if an exposure incident occurs, including the method of reporting the incident and the medical follow-up that will be made available; (L) Information on the post-exposure evaluation and follow-up that the employer is required to provide for the employee following an exposure incident; (M) An explanation of the signs and labels and/ or color coding required by paragraph (g)(1); and (N) An opportunity for interactive questions and answers with the person conducting the training session. (viii) The person conducting the training shall be knowledgeable in the subject matter covered by the elements contained in the training program as it relates to the workplace that the training will address. (ix) Additional Initial Training for Employees in HIV and HBV Laboratories and Production Facilities. Employees in HIV or HBV research laboratories and HIV or HBV production facilities shall receive the following initial training in addition to the above training requirements. (A) The employer shall assure that employees demonstrate proficiency in standard microbiological practices and techniques and in the practices and operations specific to the facility before being allowed to work with HIV or HBV. (B) The employer shall assure that employees have prior experience in the handling of human pathogens or tissue cultures before working with HIV or HBV. (C) The employer shall provide a training program to employees who have no prior experience in handling human pathogens. Initial work activities shall not include the handling of infectious agents. A progression of work activities shall be assigned as techniques are learned and proficiency is developed. The employer shall assure that employees participate in work activities involving infectious agents only after proficiency has been demonstrated. (h) Recordkeeping. (i) Medical Records. (A) The employer shall establish and maintain an accurate record for each employee with occupational exposure, in accordance with 29 CFR 1910.1020. (ii) This record shall include: (A) The name and social security number of the employee; (B) A copy of the employee's hepatitis B vaccination status including the dates of all the hepatitis B vaccinations and any medical records relative to the employee's ability to receive vaccination as required by paragraph (f)(2); (C) A copy of all results of examinations, medical testing, and follow-up procedures as required by paragraph (f)(3); (D) The employer's copy of the healthcare professional's written opinion as required by paragraph (f)(5); and (E) A copy of the information provided to the healthcare professional as required by paragraphs (f)(4)(ii)(B)(C) and (D). (iii) Confidentiality. The
employer shall ensure that employee medical records required by paragraph (h)(1) are: ..1910.1030(h)(1)(iii)(A) (A) Kept confidential; and (B) Not disclosed or reported without the employee's express written consent to any person within or outside the workplace except as required by this section or as may be required by law. (iv) The employer shall maintain the records required by paragraph (h) for at least the duration of employment plus 30 years in accordance with 29 CFR 1910.1020. (2) Training Records. (i) Training records shall include the following information: (A) The dates of the training sessions; (B) The contents or a summary of the training sessions; (C) The names and qualifications of persons conducting the training; and (D) The names and job titles of all persons attending the training sessions. (ii) Training records shall be maintained for 3 years from the date on which the training occurred. (3) Availability. (i) The employer shall ensure that all records required to be maintained by this section shall be made available upon request to the Assistant Secretary and the Director for examination and copying. ..1910.1030(h)(3)(ii) (ii) Employee training records required by this paragraph shall be provided upon request for examination and copying to employees, to employee representatives, to the Director, and to the Assistant Secretary. (iii) Employee medical records required by this paragraph shall be provided upon request for examination and copying to the subject employee, to anyone having written consent of the subject employee, to the Director, and to the Assistant Secretary in accordance with 29 CFR 1910.1020. (4) Transfer of Records. (i) The employer shall comply with the requirements involving transfer of records set forth in 29 CFR 1910.1020(h). (ii) If the employer ceases to do business and there is no successor employer to receive and retain the records for the prescribed period, the employer shall notify the Director, at least three months prior to their disposal and transmit them to the Director, if required by the Director to do so, within that three month period. (i) Dates. ..1910.1030(i)(1) (1) Effective Date. The standard shall become effective on March 6, 1992. (2) The Exposure Control Plan required by paragraph (c) of this section shall be completed on or before May 5, 1992. (3) Paragraph (g)(2) Information and Training and (h) Recordkeeping shall take effect on or before June 4, 1992. (4) Paragraphs (d)(2) Engineering and Work Practice Controls, (d)(3) Personal Protective Equipment, (d)(4) Housekeeping, (e) HIV and HBV Research Laboratories and Production Facilities, (f) Hepatitis B Vaccination and Post-Exposure Evaluation and Follow-up, and (g)(1) Labels and Signs, shall take effect July 6, 1992. [56 FR 64004, Dec. 06, 1991, as amended at 57 FR 12717, April 13, 1992; 57 FR 29206, July 1, 1992; 61 FR 5507, Feb. 13, 1996]
APPENDIX 5  Phlebotomy Checklist

Blood Collection Tray Checklist

Per Tray:

- 10 21G Butterfly needles with Luer Adapters
- 10 Alcohol Swabs
- 15 Band-Aids
- 15 Gauze pads
- 5 Vacutainer holders
- complete set of extra, unlabeled collection tubes
- 2 Tourniquets
- 1 Smelling salts
- 1 Timer or stopwatch
- 2 Pencils/ pens
- Latex gloves
- 1 Hemostats
- 1 Adhesive tape
- 1 Scissors

~10 min before draw:

- 1 styrofoam ice bath filled with ice

Per participant:

- 1 Blood tube rack with 4 draw tubes labeled and numbered.
- 1 Health ABC Phlebotomy/ Processing Form

At the Phlebotomy Station:

- Supply of disposable 21-gauge needles
- Basin
- Cold cloth
- Tube mixer
- Biohazard containers
- Needle/ Sharps container
- Paper towels
APPENDIX 6 Sample letter to physician regarding alert value

June 13, 1999

Abe Friedman, M.D.
5845 Centre Avenue
Pittsburgh, PA  15213

Dear Dr. Friedman:

On June 1, 1998, we performed a surveillance visit on your patient _______ at the Health ABC Clinic. [A complete blood count was obtained and the results of the white blood count are 16,000. (Alert values are <2,000 or >15,000.)] We will send the remainder of the results when we receive them from the Health ABC Coordinating Center.

All tests were performed for research purposes only and will be used to describe the health status of men and women in their seventies who are taking part in this study. These tests are not intended to replace any tests that might be ordered for a specific clinical indication. Although we do not suggest specific diagnosis or treatment, we hope this information is useful to you and your patient.

If you have any questions, please feel free to contact us at ________. Thank you for your support.

Sincerely,

Anne Newman, M.D., MPH
Health ABC Principal Investigator