Evidence Examination

GUIDING PRINCIPLES AND SCOPE

Specific methods to examine evidence varies by case type. Guidelines for the examination of the common types of evidence are presented in this procedure. If an analyst encounters any type of evidence not presented in this procedure, a supervisor shall be consulted for further guidance.

PROCEDURE

A. Note taking – general guidelines

Note taking and evidence documentation is the most important aspect of casework. Done improperly, it can jeopardize any analysis that follows. The notes are used to document the condition of the packaging and evidence, describe stains that may be found, present the results of presumptive and/or visual tests, support the conclusions of the report, and refresh the analyst's memory when required to testify in court. If the use of paper is required for notes, use a permanent medium such as ink—never pencil. Hard copy notes or sketches must be scanned for association to the case record in LIMS (as applicable).

1. Note taking starts with a description of the evidence packaging, including:

   a. Type of package – paper bag, manila envelope, zip-loc bag, etc.
   b. Condition of package – wet, bloody, etc.
   c. Type of seal – stapled, taped, unsealed.
   d. Identifying marks – a brief description of labels, tags, handwritten notations, etc.

   Each package must be labeled by the analyst with the evidence item identifier (see Evidence Control procedure for the numbering scheme), date, and his/her handwritten initials. Finding the marks in court is easier if the analyst always chooses the same location to put his or her marks.

2. Next is a description of the contents, the evidence itself. Specific suggestions concerning different types of evidence will be discussed later.
Discrepancies between the voucher, laboratory request form, and the items in the package must be clearly documented and a deviation must be completed within the LIMS as necessary. This includes, but is not limited to, items that were submitted, but were not included on the voucher. These items may also need to be examined. Give the item the next item number. If upon opening a package it was discovered that the description on the voucher was incorrect (for example, a tank top was submitted, but the voucher says "T-shirt"), use the correct description in your notes and subsequent analyses. Do not perpetuate the mistake.

Standardized worksheets are available with diagrams of pants, shirts, shoes, etc., to aid in documenting stain patterns. If a diagram must be hand-drawn, make sure it is large enough to allow room to document all of the stains present. It is preferable to have only one diagram per page. When complete, scan this worksheet to a .pdf format and attach to the case record within the LIMS.

The LIMS has specific worksheets for the documentation of different types of items (for example: cigarette butts, fingernails, general items, etc).

Digital photography may be substituted for diagrams. Each photograph must have a ruler visible in the frame, either a plain straight ruler or an x, y axis ruler. When the photograph is printed, the analyst must mark the photograph to highlight stains, damage, etc., and add the appropriate item or sample identifier, the analyst’s initials and date to the photograph. When complete, scan photographs to a .pdf format and attach to the case record within the LIMS. The original printout may be retained in the case file (if a hard copy exists) or discarded.

Each item of evidence must be marked by the analyst with the case number, voucher number, item number, date, and handwritten initials. Marking may be done by affixing a tag with the information or by writing directly on the item.

3. If corrections are made on hard copy examination documentation, a strike-through must be drawn through the error; and initialed and dated by the person making the changes. Additional notations, including interlineations, made on the examination documentation must also be initialed and dated. Never obliterate, including using “white-out,” any notes or entry in a worksheet.
If an error is found on the data recorded within in the LIMS, the corrections should be made in the LIMS by the appropriate level of user. These changes are tracked within the LIMS, including the date, time, and name of the user making the changes.

4. Each sample/stain that will be tested must be given a unique identifying number, clearly shown in the notes. See the “Evidence Control” procedure for the sample identification scheme. Each stain must be hand marked by the analyst. Marking may be done by affixing a tag with the information or by writing directly on the item.

For most tests, the LIMS will generate a functional report documenting the test and the results. It is the responsibility of the IA/RA to ensure that the appropriate reports are printed and inserted into the hard copy the case file.

B. Preparing for evidence examination

Before examining evidence, certain preparations should be made:

1. Review the Schedule of Analysis for analyses to be performed on the item(s) in the case. Review all the information provided in the case record. This includes the Communication Log, vouchers, requests for laboratory examination, any previous laboratory reports, and police reports. If further information or clarification is needed, obtain it before beginning analyses.

2. Plan your approach to the case. Certain items may have greater potential informational value than others, or may need to be analyzed first as an investigative aid.

3. Ensure that you are wearing the proper Personal Protective Equipment.

4. Prepare the work area. The bench must be clean and free of clutter. The LIMS cart should be sufficiently charged if on battery power. Both the bench and the LIMS cart mouse, keyboard, and cart handle should be wiped down with 10% bleach followed by 70% ethanol. The work area should then be covered with paper to prevent the loss of small particles of evidence and to prevent the cross-transfer of materials from one item to another. Change the paper when a new case is begun, between different types of evidence within a case (such as between victim’s and suspect’s belongings), between different vouchers in a case, or whenever necessary. Gloves should be changed as frequently as bench paper is changed, or whenever necessary.

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5. Make sure the necessary tools and reagents for the examination are clean and conveniently located, that there is adequate lighting available, and that note taking materials are at hand to record your observations.

C. Evidence examination – general guidelines
The examination of objects will be described in a general sense, covering a broad range of topics applicable to most items of evidence.

Record the Evidence Packaging as the initial documentation of each item.

NOTE: All cutting utensils, tweezers, etc. must be cleaned before and after each use. The recommended cleaning method is 10% bleach, and/or distilled water, and then 70% ethanol. Gloves should be changed between each item, and as needed. Lab coat should be changed after scraping an item.

1. Individual evidence packages that all relate to one case may be packaged in a mesh bag for convenience. This mesh bag should not be examined or counted as a packaging material. No documents, labels, or notes should be attached or written on the mesh bag. For the individual evidence packages, verify that outer packaging corresponds to lab request/voucher. Open the packaging. Avoid breaking existing seals when possible.

2. Remove items from packaging with care. Remember, materials of evidentiary value may adhere to the item and/or the packaging. Opening the evidence over bench paper will prevent the loss of these materials.

3. Examine one item at a time.

If it is known that an item still requires trace evidence examinations, place an additional sheet of thin (newspaper weight) paper on top of the regular paper prior to opening an item of evidence. When done examining the item, wrap it up in the thin paper and place the entire bundle back into the original packaging. Any trace evidence that was dislodged from the item must be retained within the thin paper.

4. Be certain that the previous item has been re-packaged before opening another item on the work surface.
5. If an item of evidence is found to be wet when opened, the item should be allowed to air dry. The item should not be heated or exposed to direct sunlight. If the item has become foul smelling, allow it to dry in the fume hood with the fan running. If mold is present, consult a supervisor to determine if further testing is suitable.

6. The initial evaluation of the evidence is a visual inspection. It may be necessary to use a high intensity light source, UV light source, or alternate light source during the inspection, especially if semen or saliva is suspected. Magnification may be necessary. IR light source may be utilized to help find stains on dark colored materials as well.

7. A tactile examination is sometimes helpful for locating some biological stains, notably semen stains. Using gloved fingertips, lightly brush over the surface of the object, feeling for changes in surface texture or stiffness.

8. Remove any easily visible surface debris such as hairs, fibers, wood fragments, etc. and return to the original package within a sealed coin envelope with appropriate markings indicating case number, voucher number, item number, date and initials. The location on the item of all trace evidence removed should be documented by diagram, photography, or described in the notes.

9. Perform the appropriate screening tests, such as Kastle-Meyer or Acid Phosphatase. The lot numbers of all reagents and control testing results must be documented prior to use to ensure that the reagent isn’t an expired lot within the LIMS.

10. All positive biological stains (KM positive, amylase positive and/or PSA positive) must be documented by notes, diagrams, and/or photography. Note the location of the stain, size, heaviness (soaked into fabric, surface smear, etc.). Each photograph must have a ruler visible in the frame, either a plain straight ruler or an x, y axis ruler.

   If it is apparent that there is a spatter pattern, consult a supervisor for guidance. Select appropriate stains for further testing based on any spatter analysis.

   Document whether or not the biological stains exhibit directionality, if applicable.

11. Cut, scrape, and/or swab the stain from the evidence item at the time of examination for the purpose of further testing.
When swabbing an area, the number of swabs collected must be recorded. Swabbing should only be done when cutting a stain is not practical or recommended.

12. When the examination of an item or voucher is complete (body fluid identification complete and appropriate samples/cuttings submitted for DNA testing), seal the packaging with a permanent seal. The original packaging must be sealed at all entry points. All seals must be individually initialed and dated across the tape edge. Barcodes and other agency identifiers on the outer packaging should not be covered or sealed over if possible.

If multiple items of evidence are separately packaged for a single case, these items may be collected and stored in a mesh bag. This mesh bag is used only for the sake of convenience in grouping related evidence, and should not be tagged, labeled, or have any documentation attached to the mesh bag itself. Transfer the evidence to the Evidence Unit or secure storage location for storage.

Since post-mortem items are not vouchered, transfer them to retained storage once they are ready for storage.

Each time a retained sample is removed for analysis, the chain of custody must reflect this. The retained sample package must be opened and re-sealed according to Departmental guidelines.

13. Unless there is case information to the contrary, all samples will be processed as if DNA typing is to be performed.
   a. For possible bloodstains that have tested positive with a presumptive test for blood, a portion of the stain or swab may be submitted for DNA extraction, depending on the case type.
   b. For possible semen stains that have tested positive with a presumptive test for semen, a portion of the stain or swab is submitted immediately for PSA testing.
   c. For sexual assault kit swabs with accompanying smears, a portion of the swab is submitted directly for DNA extraction if sperm are found on the smears. If no sperm are seen on the smear, perform PSA testing on the swab.
   d. For sexual assault kit swabs without accompanying smears, a portion of the swab is submitted for PSA testing.
e. For possible saliva samples, a portion of the stain or swab is submitted for amylase testing.

14. If a sample is positive for PSA or amylase, a portion of the stain or swab may be submitted for DNA extraction, as necessary.

15. To prepare samples for DNA extraction, label extraction tubes with the sample identifier and add one of the following:
   a. Blood – portion of bloodstain or swab about 3mm square, enough scrapings to give a light straw colored extract, or 3μL whole blood
   b. Semen – portion of semen stain about 5mm square, one third of a swab, or 3μL of whole semen
   c. Amylase – portion of stain about 5mm square or one third of a swab.
   d. Scrapings (of clothing items)
   e. Swab(s) of touched items

   Be mindful of the amount of scrapings and/or SDS swabs being placed in extraction tubes. Excessive amounts of substrate may hinder the extraction process.

Create the sample and schedule the appropriate extraction procedure for the sample (exemplars, bloodstains, semen stains, touched items, other evidence, or one-step). Scheduling a sample for an incorrect extraction process may lead to the subsequent results being declared inconclusive; see a supervisor if you have any questions about whether a particular sample is evidence or an exemplar.

All extraction tubes should be transferred to an extraction refrigerator.

When handling each sample:

1) Use a clean cutting surface for each sample, such as a lint-free wipe.
2) Use clean scissors for cutting each sample.
3) Use lint-free wipes or clean tube openers to open sample tubes and blood tubes.
4) If possible, the entirety of an item or sample should not be consumed during analysis. It is recommended that at least 25% of the sample be controlled.

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saved for future analysis, if needed. However, if in the opinion of the analyst (or for touched items), consumption of the sample is necessary to have the best chance to obtain results, the item or sample may be consumed; the notes must clearly state this.

16. During the normal course of examination in a limited access laboratory, evidence need not be sealed when left unattended for a short period of time (such as when the analyst takes a lunch break). However, measures must be taken to prevent the unattended evidence from coming into accidental contact with other items of evidence or personnel. For example, swabs and small clothing items should be returned to its containers, and larger items (such as bed sheets on an examination hanger) should be moved to areas of the laboratory where accidental contact by other personnel will be limited.

Direct any questions regarding what prevention measures should be taken to a supervisor prior to leaving the evidence unattended.

17. Evidence in the process of examination may not be left unattended overnight without first consulting with a supervisor. Without prior approval from a supervisor, all evidence must be properly sealed and returned to a secure storage location at the end of the day.

Under certain circumstances, the supervisor may allow evidence in the process of examination to be left unattended overnight. However, this practice is to be limited based on the necessity, and the risk of accidental contact with other items of evidence or personnel must be minimized (see Paragraph 16, above). For example, a supervisor may approve evidence to be left unattended overnight if an item of evidence is found to be wet when opened and must be air dried or dried in a hood with the fan running. However, the supervisor must ensure that all risks of accidental contact with other items of evidence or personnel are minimized.

D. Evidence examination – weapons

Weapons are frequently submitted for bloodstain or tissue examinations and/or for the recovery of DNA from skin cells, depending on the case scenario. Weapons can consist of knives, guns, bottles, baseball bats, and numerous other items.
Weapons should be thoroughly described and examined. Follow the general guidelines for note taking and evidence examination when examining any weapon.

**Beware of sharp objects that have penetrated their packaging and/or are loose inside their package and could inflict injury.**

Complete the General Packaging Worksheet as the initial documentation of each item.

Complete the General Item Examination Worksheet for each item.

1. Describe the general condition of the item, such as presence of rust or fingerprint powder. Certain weapons should be tethered within their evidence packaging. If not, a deviation should be logged.

2. Measure the physical dimensions of the item. In the case of a knife, this should include description of knife blade such as thickness, shape, cross-sectional shape, length, width, number of blades, brand names, etc. Photograph the weapon if any serology positive stains are found or if the item is being sampled for skin cells.

3. As necessary, examine under a magnifier, high intensity oblique light, infrared light, or stereomicroscope for traces of fibers, hairs, blood, or other materials of evidentiary value. All trace evidence removed should be documented in the notes using descriptions, diagrams, and/or photography.

4. Look carefully for directional spatters of blood on weapons. Discuss any directional stains with a supervisor before performing any analyses.

5. Knives, sheaths, or other weapons may be dismantled as necessary for further examination. Always photograph or diagram the intact items before dismantling.

6. All stains must be documented by notes, diagrams, and/or photography. Note the location of the stain, size, heaviness (soaked into fabric, surface smear, etc.), and any directionality of the stain pattern. Each photograph must have a ruler visible in the frame, either a straight ruler or an x, y axis ruler.

7. If the area being examined for blood appears clean, consider utilizing the sensitivity of the KM test, which is more sensitive than the human eye. Perform a global swab(s) of the area and KM test a small cutting. If the swab is KM negative, retain
the remainder of the swab in a coin envelope. If the swab is KM positive, consider consuming the swab for extraction.

8. Make every effort to avoid positive serology stains when sampling the handle of a weapon. Unless there is an indication that the suspect was bleeding, this technique will assist in isolating the desired DNA profile. In cases where the “handle” of the weapon is unknown (e.g., crowbar), treat each end separately as if it could have been the handle.

9. After examining a knife or other sharp object, package it in a safe manner (fastened and/or wrapped within the original packaging) for return to the Evidence Unit.

E. Evidence examination – clothing

Follow the general guidelines for note taking and evidence examination when examining any item of clothing.

Record the Evidence Packaging Worksheet as the initial documentation of each item.

Complete the Clothing Description or General Item Examination Worksheet for each separate clothing item.

1. Describe the color or pattern of the item of clothing, fabric type (denim, corduroy, etc.), fabric make-up (cotton, polyester, etc., from label, if present), and size (if marked on item). If an item is submitted inside-out, record this information.

2. Spread out the item of clothing, looking carefully at the front, back, and inside for any possible evidentiary material.

3. Describe the general cleanliness of the item of clothing. Note any defined soiled areas (biological and/or non-biological) on the garment, for example, knees, buttocks, or cuffs. Note whether the garment appears freshly washed or not (for example, wet or damp).

4. Describe any damage to clothing, which may have evidentiary value. For example, torn or missing buttons, torn or cut areas, damaged areas, or burned areas should be described.

5. Note the presence of any suspected stab holes or bullet holes. Diagram the location, orientation, size, and shape of any holes. Do not overlook the possibility that more than
one hole may be caused by a single stab or shot due to the folding of the fabric. When sampling a stain from the area of a suspected stab hole or bullet hole, **Do not** cut through or otherwise disturb the hole. Take a sample away from the existing hole.

6. Carefully examine any pockets, inside and out. The preferred method is to gently pat the outside of the pocket to determine if there are any contents. Tweezers may be used to turn pockets inside out. **Caution is advised when placing the hand in a pocket. An unexpected sharp object could cause serious injury.**

7. Carefully examine the waistband, lining, cuff area, and collar area. This may require turning an item inside out.

8. Shoes have many crevices, which could retain material of evidentiary value and therefore should be examined carefully. Look carefully in the groove between the sole and upper shoe. Shoes with tongues should be checked for blood, which may have fallen between the shoelaces.

   Shoes may be dismantled as necessary for further examination. Always photograph or diagram the intact items before dismantling.

9. Document stains by diagrams, description, and/or photography. Note the location of the stain, size, heaviness (soaked into fabric, surface smear, etc.), and any directionality of the stain pattern. Each photograph must have a ruler visible in the frame, either a plain straight ruler or an x, y axis ruler.

F. **Evidence examination – clothing (for skin cells)**

Clothing items that are scheduled to be examined for the DNA of the individual who wore the item should be processed using the scraping method. Follow the general guidelines for note taking and evidence examination when examining any item of clothing.

Complete the General Packaging Worksheet as the initial documentation of each item.

Complete the Clothing Description or General Item Examination Worksheet for each separate clothing item.

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After the steps described in E., do the following:

**IMPORTANT:** Do not perform this procedure near an air conditioning unit. In addition to a new lab coat and new gloves, the analyst should wear a mask/face shield and hair guard. For this technique, you must put on gloves in the following manner; latex gloves, cut-resistant gloves, then latex gloves as the final layer.

1. Make sure bench-top is covered with paper. Take another piece of bench paper and fold the edges on each of the four sides up to form a 1/2 to 3/4 inch high rim. Tape or staple the corners to maintain the raised edges. For small items the bench paper should be folded in half before doing this. This will serve as a collection device for the scrapings.

2. Use a clean unused razorblade to vigorously scrape the inside of the item, paying special attention to friction areas such as the cuffs and the neck line. Do not scrape too hard or you will produce too much lint. Make sure to cover the complete surface, if possible and appropriate. **If the item also contains biological stains, it is important not to include these areas when scraping.**

   The best way of doing this is to fold each item symmetrically, lay it down flat in the collection bin, and scrape the surface. Re-fold and repeat until the complete inside has been scraped. This procedure will produce lint that contains the skin cells; consider this lint as a carrier for the cells.

3. Collect the lint by brushing the fibers into one corner of the bench paper (use razorblade), use tweezers to transfer material into an extraction tube. If no fibers are visible, use the razorblade to scrape the bench paper surface into an extraction tube.

   The scrapings should be divided into two parts; one part goes to extraction. The remaining part is placed into an extraction tube and then packaged within an individual envelope, labeled, and returned to the original packaging.

G. **Evidence examination – touched clothing (for skin cells)**

   Clothing items that are scheduled to be examined for DNA left behind by an assailant after a physical struggle should be processed using either a swabbing or scraping method, as required.

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based on the material being examined. Follow the general guidelines for note taking and evidence examination when examining any item of clothing.

Complete the General Packaging Worksheet as the initial documentation of each item.

Complete the Clothing Description or General Item Examination Worksheet for each separate clothing item.

After the steps described in E., do the following:

**IMPORTANT:** Do not perform this procedure near an air conditioning unit. In addition to a new lab coat and new gloves, the analyst should wear a mask/face shield and hair guard. For this technique, you must put on gloves in the following manner; latex gloves, cut-resistant gloves, then latex gloves as the final layer.

1. Make sure the bench-top is covered with paper. Take another piece of bench paper and fold the edges on each of the four sides up to form a 1/2 to 3/4 inch high rim. Tape the corners to maintain the raised edges. For small items the bench paper should be folded in half before doing this. This will serve as a collection device for the scrapings.

2. Determine the substrate of the item of clothing being examined.

3. Based on the material, choose the best method to examine the item. Refer to the table below:

<table>
<thead>
<tr>
<th>Recommended method to use for various materials</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Scraping</strong></td>
</tr>
<tr>
<td>Cotton &amp; Cotton mixture</td>
</tr>
<tr>
<td>Polyester</td>
</tr>
<tr>
<td>Wool</td>
</tr>
</tbody>
</table>

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4. For swabbing, swab the entire area using irradiated SDS swabs prepared by the Quality Assurance team moistened with 0.01% SDS. Combine the swabs inside one extraction tube.

5. For material requiring scraping, scrape the entire area with a sterile blade and place the scrapings inside an extraction tube. Make sure to scrape the entire surface the assailant was purported to have had contact with. **If the item also contains biological stains, it is important not to include these areas when scraping or swabbing.**

6. After scraping the item, you may wipe the blade with an irradiated SDS swab to recover as much skin cell evidence as possible. Place the swab inside the same tube as the scrapings. Both the scrapings and the SDS swab will be extracted together as one sample.

H. Evidence examination – sexual assault kits

Follow the general guidelines for note taking and evidence examination when examining any sexual assault kit. Follow the general guidelines for clothing examination when examining any clothing items packaged in a sexual assault kit.

Complete the Evidence Packaging Worksheet as the initial documentation of each item.

Complete the Sexual Offense Evidence Collection Kit Inventory Worksheet and the Clothing Description Worksheet (for testing of underwear or related items) for further documentation of each separate clothing item.

1. Ensure that the name of the victim corresponds to the name listed on the paperwork in the case file.

2. Indicate whether each kit component is sealed, unsealed, not submitted, or present but “not used” (this may require opening of the envelope). Consecutive item numbers are assigned to only those items that are present and used (e.g., 1.1, 1.2, 1.3.1-1.3.2 for swab and smear pairs).
PM kits (all items packaged together): Inventory kit. Label used envelopes with an item number (see above) and the FB number (label as 1.1, 1.2, etc), analyst’s initials, and date of examination. All the envelopes, whether used or unused should contain the analyst’s initials and the identifying case number. All envelopes and any paperwork associated with the PM kit will be retained in the kit box. For PM SAKs use the Sexual Offense Evidence Collection Kit Inventory Worksheet.

PM swabs (items packaged separately): Complete the Packaging and Swab Examination Worksheet. These swabs should already have item numbers. Refer to the LIMS Evidence Manual.

Vouchered kits: Inventory kit. Label used envelopes with an item number (see above) and the FB number, analyst’s initials, and date of examination. All the envelopes, whether used or unused should contain the analyst’s initials and the identifying case number. See following for testing of the vouchered kit.

3. Underwear or related items contained within kit:

If underwear or related items (e.g., pantiliner) are in the kit, complete the Clothing Description or General Item Examination Worksheet. If stains are observed, underwear can be documented using the diagrams that are available or by a quick sketch. Photography is not generally needed.

Testing of underwear or small clothing items contained within kit:

Visually check underwear for any biological stains. Additionally, observe the underwear using an alternate light source. If any fluorescing areas are observed, circle for further testing.

If a whitish, yellowish, or fluorescing stain is observed on the underwear, test the stain with AP reagent. If the stain is AP positive, make a small cutting and submit the stain for PSA confirmation testing.

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If a pink to reddish-brown stain is observed on the underwear, test the stain with KM reagent and AP reagent, even if the area does not appear to fluoresce with the aid of the alternate light source. The possible presence of blood can mask fluorescence. Regardless of KM results, the stain needs to be AP tested. KM positive stains should be documented.

If the time since date of occurrence is more than 3 months, the analyst should submit a small portion of the area for PSA testing.

At this point, be sure that any AP positive stains submitted to PSA testing are designated a stain number. A stain number should also be designated for KM positive stains. KM positive only stains do not require further testing.

If oral sodomy is suspected or it is unclear what type of sexual contact occurred in the case, it may be necessary to send stains for amylase testing. Consult with exam supervisor as needed.

If there are no biological stains on the item(s), a diagram is not necessary; write a short description of the item.

Testing of gauze within the kit:

Examination of gauze is similar to underwear, however all AP positive and negative stains should be tested for the presumptive presence of amylase.

Note the location from which the gauze was collected. If the location from which the gauze was taken is known, this information must be included on the SAK inventory.

4. The trace evidence envelope is used by hospital personnel to collect trace evidence from the victim’s body and/or the clothing. The victim disrobes over examination paper, and the examination paper is collected.
Since trace evidence examinations are not performed in the Department of Forensic Biology, there is generally no need to examine the contents. However, if the envelope appears to contain something other than trace evidence, or markings on the envelope indicate that something other than trace evidence is present, the envelope should be opened to confirm the contents and examination should proceed if needed. If the contents of the envelope are found to be the examination paper, no further examination is needed.

5. The debris envelope is used by hospital personnel to collect loose, obvious foreign material from the victim’s body and/or the clothing.

If a debris envelope was used, note the location from which the debris was collected, or note that the location was not given. Since trace evidence examinations are not performed in the Department of Forensic Biology, there is generally no need to examine the contents.

6. The dried secretions swabs are used to collect possible biological fluids from areas other than the body cavities.

If dried secretions were taken, note the number of swabs and the location from which the secretions were collected, or note that the location was not given. Each swab must be individually labeled (1.4.1, 1.4.2). See below for further testing procedures.

**Testing of dried secretions swabs:**

Make a cutting from each of the swabs present for PSA testing. If the location from which the dried secretions swabs were taken is known, and is not from the mouth, near the mouth, anal cavity, or near the anal cavity, the swab should also be tested for amylase. Swabs from these locations are not tested for amylase. If the location is unknown, make a cutting from each swab for both PSA and amylase testing. As every case is different, please consult with a supervisor if there is something in the case description that suggests further testing is required.
7. The **fingernail scrapings (or clippings)** are used to collect trace evidence from the fingernails.

Since trace evidence examinations are not performed in the Department of Forensic Biology, there is generally no need to examine the contents. However, requests are occasionally made to examine the fingernail scrapings; discuss with a supervisor before starting any examinations of fingernail evidence. If fingernail examination has been approved, refer to Section O of this manual.

8. If a **liquid blood exemplar** is present, consult with a supervisor to make a bloodstain card. Fill out a blank stain card (FB number, victim’s name, date, and initials), insert into a Kapak envelope and seal it. The FB number should be written on the Kapak and the analyst’s initials and date of examination should be written across the seal. This may be used as an exemplar if no buccal specimen is present within the kit.

9. If a **dried blood control** is present, it is only used if there is no buccal specimen present in the kit. If it must be used, fill out a blank stain card (FB number, victim’s name, date, and initials), attach the dried blood control to it, insert into a Kapak envelope and seal it. The FB number should be written on the Kapak and the analyst’s initials and date of examination should be written across the seal.

10. The **buccal specimen** is used as the victim’s exemplar. If present, the buccal specimen would be the first choice in order to avoid potential inhibition of PCR by heme degradation products.

If no victim’s exemplar is present, and there are no serology negative body cavity swabs, it may be necessary at a later time for a supervisor to make a phone call to request one.

11. The **pulled head hair and pulled pubic hair** are collected as exemplars for any future microscopic hair comparisons.

Since trace evidence examinations are not performed in the Department of Forensic Biology, there is generally no need to examine the contents. However, requests are occasionally made to use the pulled head hair for exemplar DNA testing; generally, hair DNA testing is not performed until hair comparisons have been made by the NYPD forensic laboratory.

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12. The **pubic hair combings** are used to collect possible trace evidence from the pubic hair of the victim.

Since trace evidence examinations are not performed in the Department of Forensic Biology, there is generally no need to examine the contents.

13. The “**body cavity**” **swabs** (oral, perianal, anal, vulvar, vaginal/penile, and cervical) are used to collect possible biological fluids from those areas; the smears are used for a sperm search.

**Testing of body cavity swabs:**

Visually check the swabs for the presence of biological fluids. If the swabs are pink to reddish-brown in color, test with KM reagent; note the results.

Refer to the Sexual Assault Kit Processing Flow Charts for guidance.

Stain one smear accompanying each set of body cavity swabs using the Christmas Tree staining procedure (refer to Christmas Tree Stain for Spermatozoa section in the Forensic Biology Serology Manual) and examined for the presence of sperm; this search need not be exhaustive and should take no longer than five minutes (per smear). It is not necessary to estimate the number of sperm present, but the relative amount (one sperm head, numerous sperm heads, etc.) may be noted.

If sperm is found on a smear, make a cutting from each positive location on relevant swabs (vulvar, penile, scrotal) for amylase testing.

If no sperm is found on a smear, make a cutting from each negative location for PSA confirmatory testing. Pertinent swabs (vulvar, penile and scrotal) must also be tested for the presence of amylase.

**Body cavity swabs** (vaginal, cervical, oral, and anal) **should not be tested for the presence of amylase.** Swabs labeled “perianal/anal” **should not routinely be tested for amylase; however, they may be tested if clearly marked as “perianal”**.

14. Return all swabs and smears to their respective envelopes.
15. Questionnaires, forms, and body diagram sheets are intended for the use of the medical personnel. Any such paperwork found in the kit that is filled-out with handwritten information should be copied for retention with the case record—as a physical copy in the case file and a .pdf attachment in LIMS (as applicable); leave all originals in the kit. If present, these filled-out documents are considered administrative records and must be labeled with the Forensic Biology case number. No item number designation is needed if present.

16. Photographs and/or other paperwork are not supposed to be included in a kit. If present, make a note of it; leave them in the kit. No item number is assigned if present.

17. After kit examination is complete, the kit is now ready to be closed. After the kit is closed, the kit should be placed in a secure location.

**Closing of negative kits:**

If the kit is negative for semen and amylase, and there is no other evidence to examine, the case is finished and should be submitted to Quality Assurance for reanalysis consideration.

If a buccal specimen is present, the analyst should place the swab(s) in a coin envelope labeled with the FB number, voucher number, item number, victim name, analyst’s initials, and date of examination. The coin envelope should be placed in a Kapak envelope and heat sealed. The FB number should be written on the Kapak and the analyst’s initials and date of examination should be written across the seal. The buccal should be transferred to a storage location.

If no buccal specimen was present in the kit, retain serology negative body cavity swabs to be used as an exemplar. The oral swab is the preferable choice to be used as an exemplar in the absence of a true exemplar.

Each envelope within the kit should be sealed with evidence tape. The entire vouchered kit or the post mortem items (PM kit) kit can be placed in a secure storage location.

If the kit is negative for semen and amylase, and there is additional evidence, a supervisor will determine whether or not the evidence needs to be signed in and examined.

**Closing of positive kits:**

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If positive swabs/stains/smears were found, see below for guidelines on the cutting of samples for extraction.

If there is additional evidence, a supervisor will determine whether or not the evidence needs to be signed in and examined.

**Dried secretions swabs**

Whether or not a dried secretions swab continues on for DNA extraction, and if so, which type of DNA extraction, depends on a number of factors: location the sample was taken from, nature of the body fluid present, presence or absence of a suspect, and what other swabs or other evidence has been submitted in the case. Refer to the Sexual Assault Kit Processing Flowcharts for guidance.

If PSA positive, make a second cutting from one swab **from each listed location** that is positive for differential extraction. If the location from which the swabs were taken is unknown, make a cutting of one swab from each separate packaging to go on for differential extraction.

If a swab is PSA negative and amylase positive, the decision on further testing depends on the locations the swab was taken from (if known) and whether the case has a suspect.

PSA negative, amylase positive dried secretions from external areas should be sent to extraction. In addition, a supervisor may need to make a phone call to determine the status of the case.

**Body cavity swabs (oral, perianal, anal, vulvar, vaginal/penile, cervical)**

If sperm is found on a smear, a cutting from the accompanying swab can go for extraction. **If sperm is found on a perianal/anal smear, cuttings from both swabs are combined for extraction.** If multiple smears are sperm positive from similar areas, it is not necessary to cut all swabs for DNA extraction. For the purposes of sending samples onto extraction, vaginal swabs should be sent first, then cervical swabs, then vulvar swabs. Therefore, if all three swabs are sperm search positive, only send the vaginal swab for extraction. Refer to the Sexual Assault Kit Processing Flowcharts for guidance.

If a swab is PSA positive, a cutting from the swab can go for extraction. If multiple swabs are PSA positive from similar areas, it is not necessary to cut all swabs for DNA extraction. Refer to the Sexual Assault Kit Processing Flowcharts for guidance.
If a vulvar swab is sperm/PSA negative but amylase positive, check to see if the case has a named suspect (listed first and last name). If so, make a cutting from one swab that is amylase positive and submit this cutting to an appropriate DNA extraction. If multiple swabs are amylase positive, it is not necessary to cut all swabs for DNA extraction. Refer to the Sexual Assault Kit Processing Flowcharts for guidance. If the case has no named suspect, consult with a supervisor. It may be necessary for the supervisor to make a phone call to determine the status of the case.

If a penile swab is sperm/PSA negative but amylase positive, make a cutting from the swab and submit to the appropriate DNA extraction.

**Underwear or small items**

For PSA positive stains, cut positive stain(s) for differential extraction. For multiple suspects, it may be necessary to send multiple stains. Consult a supervisor.

In the event that there are amylase positive stains, the decision for further testing is case dependent. Consult a supervisor.

If a buccal specimen is present, make a cutting for extraction. Following this, place the remainder of the swab(s) in a coin envelope labeled with the FB number, voucher number, item number, victim name, analyst’s initials, and date of examination. The coin envelope should be placed in a Kapak envelope and heat sealed. The FB number should be written on the Kapak and the analyst’s initials and date of examination should be written across the seal.

The exemplar should be placed in a secure storage location.

After cutting all positive items, each envelope within the kit should be sealed with evidence tape and returned to the kit. Seal the kit and return to a secure storage location.
Sexual Assault kit processing flow chart

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Sexual Assault kit processing flow chart

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Sexual Assault kit processing flow chart

- Stains smear and examine for sperm
  - Sperm Positive?
    - No: Cut one swab for PSA testing
    - Yes: Cut one sperm positive swab for differential extraction
      - PSA Positive?
        - No: Done with items - Return to kit
          - Yes: Proceed with (vi) exemplar
  - Oral Swabs:
    - Cut one swab for (vi) exemplar
      - PSA Positive?
        - No: Done with items - Return to kit
          - Yes: Serology Report

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Sexual Assault kit processing flow chart

**Vulvar Swabs:**

1. Stain smear and examine for sperm
   - Sperm Positive? Yes: Cut one sperm positive swab for differential extraction; Test swab for amylase
   - Sperm Positive? No: Cut one sperm positive swab for PSA and amylase testing
   - PSA Positive? Yes: Cut one PSA-positive swab for differential extraction; Test swab for amylase
   - PSA Positive? No: Amylase Positive? Yes: Is there a suspect? Yes: Process (v) exemplar; Done with items - Return to Kit
   - PSA Positive? No: Amylase Positive? No: Cut one amylase positive swab that is semen negative for extraction for Y-testing
   - Amylase Positive? No: Serology Report

*Cut for extraction if vaginal and cervical swabs are semen negative or if case has multiple suspects.
**Cut for Y testing if no semen positive samples in the case
^Suspect must have a first and last name listed
Sexual Assault kit processing flow chart

1. Stain smear and examine for sperm
2. Sperm Positive?
   - Yes: Cutone swab for differential extraction
   - No: Cut one swab for PSA and amylase testing
3. Cutone swab for PSA and amylase testing
4. PSA Positive?
   - Yes: Cutone swab for differential extraction
   - No: Cut one swab for robotic extraction
5. Cutone swab for differential extraction
6. Cutone swab for robotic extraction
7. Proceed (v) exemplar
8. Done with items - Return to kit
9. Done with items - Return to kit
10. Done with items - Return to kit

Penile Swabs:
Sexual Assault kit processing flow chart

**Vaginal, Cervical, Perianal, and Anal Swabs:**

- Stain smear and examine for sperm
- Sperm Positive? Yes → PSA Positive? Yes → Serology Report
- Sperm Positive? No → Cut one sperm positive swab from each area for differential extraction
- Cut one sperm positive swab from each area for differential extraction
- Process (s) example
- Done with items - Return to kit
- Done with items - Return to Kit
- PSA Positive? No → Cut one of each swab for PSA testing
- PSA Positive? Yes → Serology Report

*If multiple suspects are involved, discuss case with exam supervisor.

+ For sperm positive unlabeled perianal/anal smears, combine a portion of each swab for differential extraction.

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Sexual Assault kit processing flow chart

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I. Evidence examination – male suspect kits

Although testing procedures are similar to sexual assault kit examination, the goal is to try to find victim DNA when examining any suspect kit. This should be kept in mind during examination of all items within the suspect kit.

In addition to this manual, follow the general guidelines for note taking and evidence examination, and the guidelines for clothing examination when examining any clothing items.

Use an Evidence Packaging Worksheet for initial documentation of each suspect kit.

Use the Suspect Evidence Collection Kit Inventory and Clothing Description Worksheet (for testing of underwear or related items) forms for further documentation.

1. Note the name of the suspect and information about when and where the kit was collected. Ensure that the name of the suspect corresponds to the name listed on the paperwork in the case file.

2. Inventory kit: The LIMS will assign an item number to each used envelope. Affix a LIMS packaging label to each envelope. The analyst must mark all envelopes with their initials and date of examination.

As prompted by the Suspect Evidence Collection Kit Inventory, indicate whether each kit component is sealed, unsealed, not submitted, or present but “not used” (this may require opening of the envelope).

If a buccal specimen or other exemplar sample is contained within the kit, contact a supervisor immediately to create a suspect file. Suspect file creation is only necessary if an exemplar sample is present.

Suspect file creation:

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A supervisor is responsible for creating the suspect file. The supervisor must:

- Create a LIMS record and Schedule of Analysis
- Include the following paperwork in the file upon completion of kit examination:
  - 61 form (NYPD complaint report)
  - Original request for laboratory examination forms
  - Evidence voucher
  - Evidence packaging worksheet
  - Completed kit inventory worksheet

After creation of a suspect file, the buccal swab is cut and duplicate cut for extraction in accordance with laboratory guidelines.

3. **Underwear or related items contained within kit:**

If **underwear or related items** are in the kit, examine them using the Clothing Description Worksheet.

**Testing of underwear or small clothing items contained within kit:**

Visually check underwear for any biological stains. Then observe the underwear using an alternate light source. If any fluorescing areas are observed, circle for further testing.

If a potentially biological or fluorescing stain is observed on the underwear, test the stain with AP reagent. If the stain is AP positive, make a small cutting for PSA and amylase testing. If the stain is AP negative, make a small cutting for amylase testing.

If a pink to reddish-brown stain is observed on the underwear, test the stain with KM reagent as well as AP reagent, even if the area does not appear to fluoresce with the aid of the alternate light source. Regardless of KM results, the stain needs to be AP tested. If the stain is AP positive, make a small cutting for PSA and amylase testing. If the stain is AP negative, make a small cutting and submit the stain for amylase testing. KM positive stains should be documented but do not require further testing.

In any situation, if the stain is AP negative and the time from the date of occurrence to the date of kit examination is more than 3 months, make a cutting of the stain for PSA testing.
testing to confirm negative results (for semen samples older than 3 months, AP can degrade and thus testing may yield a negative AP result).

**Remember that the goal is to try to find victim DNA.** Therefore, non-fluorescing stains may need to be tested further. Refer to the Suspect Kit Processing Flow Charts for guidance. Stain location and the case scenario will determine what stains need further testing. As every case is different, consult with a supervisor as needed.

At this point, be sure that any stains submitted to PSA and/or amylase testing and KM positive stains are designated a stain number/letter. Only KM, PSA, or amylase positive stains should be diagrammed.

If there are no biological stains on the item(s), a diagram is not necessary; write a short description of the item using a Clothing Description Worksheet.

4. **The debris envelope** is used by hospital personnel to collect loose, obvious foreign material from the victim’s body and/or the clothing.

If a debris envelope was used, note the location from which the debris was collected, or note that the location was not given. Trace evidence examinations are not performed in the Department of Forensic Biology.

5. **The dried secretions swabs** are used to collect possible biological fluids from areas other than the body cavities. This could include, for example, semen from the skin or saliva from bite marks.

If dried secretions were taken, note the number of swabs and the location from which the secretions were collected, or note that the location was not given. Each swab must be individually itemized.

**Testing of dried secretions swabs:**

Visually check the swabs for the presence of biological fluids. If the swabs are pinkish to reddish-brown in color, test with KM reagent, and note the results.

Make a cutting from each of the swabs present for PSA testing. If the location from which the dried secretions swabs were taken is known, and is not from the mouth, near Controlled versions of Department of Forensic Biology Manuals only exist in the Forensic Biology Qualtrax software. All printed versions are non-controlled copies.

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the mouth, anal cavity, or near the anal cavity, the swab should also be tested for amylase. Swabs from these locations are not tested for amylase. If the location is unknown, make a cutting from each swab for both PSA and amylase testing. As every case is different, please consult with a supervisor if there is something in the case description that suggests further testing is required.

6. The **fingernail scrapings (or clippings)** are used to collect trace evidence from the fingernails.

Trace evidence examinations are not performed in the Department of Forensic Biology. However, requests are occasionally made to examine the fingernail scrapings; discuss with a supervisor before starting any examinations of fingernail evidence. If fingernail examination has been approved, refer to Section O of this manual.

7. The **chest hair combings** are used to collect possible trace evidence from the chest hair of the suspect.

Trace evidence examinations are not performed in the Department of Forensic Biology.

8. The **oral swabs** are used to collect possible biological fluids from that area; the smears are used for a sperm search in cases with a male victim.

Refer to the Suspect Kit Processing Flow Charts for guidance.

For male victims:

Visually check the swabs for the presence of biological fluids. If the swabs are pink to reddish-brown in color, test with KM reagent; note the results.

Stain one smear accompanying each set of body cavity swabs using the Christmas Tree staining procedure (refer to Christmas Tree Stain for Spermatozoa section in the Forensic Biology Serology Manual) and examine for the presence of sperm; this search need not be exhaustive and should take no longer than five minutes (per smear). It is not necessary to estimate the number of sperm present, but the relative amount (one sperm head, numerous sperm heads, etc.) may be noted.

If no sperm is found on a smear, make a cutting for PSA testing.
For female victims:

In most cases, oral swabs and smears should not be tested. As every case is different, please consult with a supervisor if there is something in the case description that suggests further testing is required.

9. The **pulled head hair and pulled pubic hair** are collected as exemplars for any future microscopic hair comparisons.

Trace evidence examinations are not performed in the Department of Forensic Biology. However, requests are occasionally made to use the pulled head hair for exemplar DNA testing.

10. The **facial hair combings and pubic hair combings** are used to collect possible trace evidence from the facial hair and pubic hair of the suspect.

Trace evidence examinations are not performed in the Department of Forensic Biology.

11. The **penile and scrotal swabs** are used to collect possible biological fluids from those areas; the smears are used for a sperm search.

**Testing of penile and scrotal swabs:**

Visually check the swabs for the presence of biological fluids. If the swabs are pink to reddish-brown in color, test with KM reagent; note the results.

Refer to the Suspect Kit Processing Flow Charts for guidance.

For male or female victims:

Stain one smear accompanying each set of body cavity swabs using the Christmas Tree staining procedure (refer to Christmas Tree Stain for Spermatozoa section in the Forensic Biology Serology Manual) and examine for the presence of sperm; this search need not be exhaustive and should take no longer than five minutes (per smear). It is not necessary to estimate the number of sperm present, but the relative amount (one sperm head, numerous sperm heads, etc.) may be noted.
If sperm is found on a smear, make a cutting from each positive location for amylase testing.

If no sperm is found on a smear, make a cutting from each negative location for PSA and amylase testing.

12. The anal swabs are used to collect possible biological fluids from that area; the smears are used for a sperm search.

Refer to the Suspect Kit Processing Flow Charts for guidance.

For male victims:

Visually check the swabs for the presence of biological fluids. If the swabs are pink to reddish-brown in color, test with KM reagent; note the results.

Stain one smear accompanying each set of body cavity swabs using the Christmas Tree staining procedure (refer to Christmas Tree Stain for Spermatozoa section in the Forensic Biology Serology Manual) and examine for the presence of sperm; this search need not be exhaustive and should take no longer than five minutes (per smear). It is not necessary to estimate the number of sperm present, but the relative amount (one sperm head, numerous sperm heads, etc.) may be noted.

If no sperm is found on a smear, make a cutting for PSA testing.

For female victims:

In most cases, anal swabs and smears should not be tested. As every case is different, please consult with a supervisor if there is something in the case description that suggests further testing is required.

13. The buccal specimen is used as the suspect’s exemplar. If present, the buccal specimen would be the first choice in order to avoid potential inhibition of PCR by heme-degradation products.
14. The questionnaires, body diagram sheets, and instruction sheets are intended for the use of the medical personnel. If present, make a copy only of the questionnaire and body diagram sheets for retention with the case record—as a physical copy in the case file and a .pdf attachment in LIMS (as applicable); leave all originals in the kit. No item number is assigned if present. Label each page with the suspect file number, voucher number, analyst’s initials, and date of examination.

15. Photographs are not supposed to be included in a kit. If present, make a note of it, alert a supervisor, and leave them in the kit. Label with FB number, date of examination, and analyst’s initials. No item number is assigned if present.

16. If no positive swabs/stains/smears were found, make cuttings of appropriate swabs or stains as necessary. Refer to the Suspect Kit Processing Flow Charts for guidance.

17. If positive swabs/stains/smears were found, see below for guidelines on the cutting of samples for extraction. Refer to the Suspect Kit Processing Flow Charts for guidance.

The following kit closing information is for both female and male victims. Use the pertinent information for each case.

Underwear

PSA positive stains should be sent for differential extraction.

PSA negative stains that are KM and/or amylase positive should be sent for robotic extraction.

Dried secretion swabs

If PSA positive, make a cutting from one swab from each listed location that is positive for differential extraction. If the location from which the swabs were taken is unknown, make a cutting from one swab to go on for a differential extraction.

If a swab is KM positive and PSA negative, make a cutting from one swab from each listed location that is KM positive for robotic extraction.

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If a swab is amylase positive, and PSA and KM negative, refer to the Suspect Kit Processing Flow Charts for guidance.

If a swab is PSA and amylase negative, refer to the Suspect Kit Processing Flow Charts for guidance.

**Penile and scrotal swabs**

If a swab is sperm/PSA positive, make a cutting from each positive location for differential extraction.

sperm/PSA negative stains that are KM and/or amylase positive should be sent for robotic extraction.

If a swab is sperm/PSA and amylase negative, the decision on further testing depends on the sex of the victim. Refer to the Suspect Kit Processing Flow Charts for guidance.

**Oral and anal swabs**

If a swab is sperm/PSA positive, make a cutting from each positive location for differential extraction.

If a swab is sperm/PSA negative, consult with a supervisor.

18. After cutting all pertinent items, return all swabs and smears to their respective envelopes. Seal all kit envelopes with evidence tape and return to the kit. Seal the kit and return to a secure storage location.

**J. Evidence examination – female suspect kits**

Although testing procedures are similar to sexual assault kit examination, the goal is to try to find victim DNA when examining any suspect kit. This should be kept in mind during examination of all items within the suspect kit.

In addition to this manual, follow the general guidelines for note taking and evidence examination, and the guidelines for clothing examination when examining any clothing items. Controlled versions of Department of Forensic Biology Manuals only exist in the Forensic Biology Qualtrax software. All printed versions are non-controlled copies.
Use an Evidence Packaging Worksheet for initial documentation of each suspect kit.

Use the Suspect Evidence Collection Kit Inventory and Clothing Description Worksheet (for testing of underwear or related items) forms for further documentation.

1. Note the name of the suspect and information about when and where the kit was collected. Ensure that the name of the suspect corresponds to the name listed on the paperwork in the case file.

2. **Inventory kit:** The LIMS will assign an item number to each used envelope. Affix a LIMS packaging label to each envelope. The analyst must mark all envelopes with their initials and date of examination.

As prompted by the Suspect Evidence Collection Kit Inventory, indicate whether each kit component is sealed, unsealed, not submitted, or present but “not used” (this may require opening of the envelope).

**If a buccal specimen or other exemplar sample is contained within the kit, contact a supervisor immediately to create a suspect file. Suspect file creation is only necessary if an exemplar sample is present.**

**Suspect file creation:**

A supervisor is responsible for creating the suspect file. The supervisor must:

- Create a LIMS record and Schedule of Analysis
- Include the following paperwork in the file upon completion of kit examination:
  - 61 form (NYPD complaint report)
  - Original request for laboratory examination forms
  - Evidence voucher
  - Evidence packaging worksheet
  - Completed kit inventory worksheet

After creation of a suspect file, the buccal swab is cut and duplicate cut for extraction in accordance with laboratory guidelines.
3. **Underwear or related items contained within kit:**

If **underwear or related items** are in the kit, examine them using the Clothing Description Worksheet. If stains are observed, underwear can be documented using the diagrams that are available or by a quick sketch. Photography is not generally needed.

**Testing of underwear or small clothing items contained within kit:**

**For male victims:**

Visually check underwear for any biological stains. Additionally, observe the underwear using an alternate light source. If any fluorescing areas are observed, circle for further testing.

If a potentially biological or fluorescing stain is observed on the underwear, test the stain with AP reagent. If the stain is AP positive, make a small cutting for PSA and amylase testing. If the stain is AP negative, make a small cutting for amylase testing.

If a pink to reddish-brown stain is observed on the underwear, test the stain with KM reagent and AP reagent, even if the area does not appear to fluoresce with the aid of the alternate light source. Regardless of KM results, the stain needs to be AP tested. If the stain is AP positive, make a small cutting for PSA and amylase testing. If the stain is AP negative, make a small cutting for amylase testing. KM positive stains should be documented.

In any situation, if the stain is AP negative and the time from the date of occurrence to the date of kit examination is more than 3 months, the analyst should make a cutting of the area for PSA testing to confirm negative results (for semen samples older than 3 months, AP can degrade and thus testing may yield a negative AP result).

At this point, be sure that any stains submitted to PSA and/or amylase testing and KM positive stains are designated a stain number/letter.

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If there are no biological stains on the item(s), a diagram is not necessary; write a short description of the item using a Clothing Description Worksheet.

**For female victims:**

Visually check underwear for any biological stains. Additionally, observe the underwear using an alternate light source. If any fluorescing areas are observed, circle for further testing.

If a fluorescing stain is observed on the underwear, make a small cutting for amylase testing. Designate a stain number/letter to each fluorescing area.

If a pink to reddish-brown stain is observed on the underwear, test the stain with KM reagent. Consult with a supervisor.

**Remember that the goal is to try to find victim DNA.** Therefore, non-fluorescing stains may need to be tested further. Refer to the Suspect Kit Processing Flow Charts for guidance. Stain location and the case scenario will determine what stains need further testing. Consult with a supervisor as needed.

At this point, be sure that any KM, PSA, or amylase positive stains are designated a stain number/letter.

If there are no biological stains on the item(s), a diagram is not necessary; write a short description of the item using a Clothing Description Worksheet.

4. The **debris envelope** is used by hospital personnel to collect loose, obvious foreign material from the victim’s body and/or the clothing.

If a debris envelope was used, note the location from which the debris was collected, or note that the location was not given. Trace evidence examinations are not performed in the Department of Forensic Biology.

5. The **dried secretions swabs** are used to collect possible biological fluids from areas other than the body cavities. This could include semen from the skin or saliva from bite marks, for example.

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If dried secretions were taken, note the number of swabs and the location from which the secretions were collected, or note that the location was not given. Each swab must be individually itemized.

Refer to the Suspect Kit Processing Flow Charts for guidance.

**For male victims:**

Visually check the swabs for the presence of biological fluids. If the swabs are pinkish to reddish-brown in color, test with KM reagent, and note the results. Make a cutting from each of the swabs present for PSA testing. If the location from which the dried secretions swabs were taken is known, and is not from the mouth, near the mouth, anal cavity, or near the anal cavity, the swab should also be tested for amylase. Swabs from these locations are not typically tested for amylase. If the location is unknown, make a cutting from each swab for both PSA and amylase testing. As every case is different, please consult with a supervisor if there is something in the case description that suggests further testing is required.

**For female victims:**

Visually check the swabs for the presence of biological fluids. If the swabs are pinkish to reddish-brown in color, test with KM reagent, and note the results. Make a cutting from each of the swabs present for amylase testing. Swabs with locations from the mouth, near the mouth, anal cavity, or near the anal cavity should not automatically go on for amylase testing. As every case is different, please consult with a supervisor if there is something in the case description that suggests further testing is required.

6. The **fingernail scrapings (or clippings)** are used to collect trace evidence from the fingernails.

Trace evidence examinations are not performed in the Department of Forensic Biology. However, requests are occasionally made to examine the fingernail scrapings; discuss with a supervisor before starting any examinations of fingernail evidence. If fingernail examination has been approved, refer to Section O of this manual.
7. The *chest hair combings* are used to collect possible trace evidence from the chest hair of the suspect.

Trace evidence examinations are not performed in the Department of Forensic Biology.

8. The **oral swabs** are used to collect possible biological fluids from that area; the smears are used for a sperm search.

Refer to the Suspect Kit Processing Flow Charts for guidance.

**For male victims:**

Visually check the swabs for the presence of biological fluids. If the swabs are pinkish to reddish-brown in color, test with KM reagent; note the results.

Stain one smear accompanying each set of body cavity swabs using the Christmas Tree staining procedure (refer to Christmas Tree Stain for Spermatozoa section in the Forensic Biology Serology Manual) and examine for the presence of sperm; this search need not be exhaustive and should take no longer than five minutes (per smear). It is not necessary to estimate the number of sperm present, but the relative amount (one sperm head, numerous sperm heads, etc.) may be noted.

If no sperm is found on a smear, make a cutting for PSA testing.

**For female victims:**

In most cases, oral swabs and smears should not be tested. As every case is different, please consult with a supervisor if there is something in the case description that suggests further testing is required.

9. The **pulled head hair and pulled pubic hair** are collected as exemplars for any future microscopic hair comparisons.

Trace evidence examinations are not performed in the Department of Forensic Biology. However, requests are occasionally made to use the pulled head hair for exemplar DNA testing; generally, hair DNA testing is not performed until hair comparisons have been made by the NYPD forensic laboratory.

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10. The **facial hair combings and pubic hair combings** are used to collect possible trace evidence from the facial hair and pubic hair of the suspect.

Trace evidence examinations are not performed in the Department of Forensic Biology.

11. The **vaginal and cervical swabs** are used to collect possible biological fluids from those areas; the smears are used for a sperm search.

**Testing of vaginal and cervical swabs:**

Visually check the swabs for the presence of biological fluids. If the swabs are pinkish to reddish-brown in color, test with KM reagent; note the results.

Refer to the Suspect Kit Processing Flow Charts for guidance.

**For male victims:**

Stain one smear accompanying each set of body cavity swabs using the Christmas Tree staining procedure (refer to Christmas Tree Stain for Spermatozoa section in the Forensic Biology Serology Manual) and examine for the presence of sperm; this search need not be exhaustive and should take no longer than five minutes (per smear). It is not necessary to estimate the number of sperm present, but the relative amount (one sperm head, numerous sperm heads, etc.) may be noted.

If no sperm is found on a smear, make a cutting from each negative location for PSA testing.

**For female victims:**

In most cases, vaginal and cervical swabs and smears should not be tested. As every case is different, please consult with a supervisor if there is something in the case description that suggests further testing is required.

12. The **anal swabs** are used to collect possible biological fluids from those areas; the smears are used for a sperm search.
For male victims:

Visually check the swabs for the presence of biological fluids. If the swabs are pinkish to reddish-brown in color, test with KM reagent; note the results.

Stain one smear accompanying each set of body cavity swabs using the Christmas Tree staining procedure (refer to Christmas Tree Stain for Spermatozoa section in the Forensic Biology Serology Manual) and examine for the presence of sperm; this search need not be exhaustive and should take no longer than five minutes (per smear). It is not necessary to estimate the number of sperm present, but the relative amount (one sperm head, numerous sperm heads, etc.) may be noted.
If no sperm is found on a smear, make a cutting for PSA testing.

For female victims:

In most cases, anal swabs and smears should not be tested. As every case is different, please consult with a supervisor if there is something in the case description that suggests further testing is required.

13. The buccal specimen is used as the suspect’s exemplar. If present, the buccal specimen would be the first choice in order to avoid potential inhibition of PCR by heme-degradation products.

14. The questionnaire, body diagram sheets, and instruction sheets are intended for the use of the medical personnel. If present, make a copy only of the questionnaire and body diagram sheets for retention with the case record—as a physical copy in the case file and a .pdf attachment in LIMS (as applicable); leave all originals in the kit. No item number is assigned if present. Label each page with the suspect file number, voucher number, analyst’s initials, and date of examination.

15. Photographs are not supposed to be included in a kit. If present, make a note of it, alert a supervisor, and leave them in the kit. Label with FB number, date of examination, and analyst’s initials. No item number is assigned if present.

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16. If no positive swabs/stains/smears were found, make cuttings of appropriate swabs or stains as necessary. Refer to the Suspect Kit Processing Flow Charts for guidance.

17. If positive swabs/stains/smears were found, see below for guidelines on the cutting of samples for extraction.

Refer to the Suspect Kit Processing Flow Charts for guidance.

Underwear

PSA positive stains should be sent for differential extraction.
Amylase positive, semen negative stains should be sent for other extraction.

If a stain is KM positive, consult with a supervisor.

If a stain is PSA and amylase negative, consult with a supervisor.

Dried secretion swabs

If PSA positive, make a second cutting from one swab **from each listed location** that is positive for differential extraction. If the location from which the swabs were taken is unknown, make a cutting from one swab to go on for a differential extraction.

If a swab is KM positive and PSA negative, make a cutting from one swab **from each listed location** that is KM positive for blood extraction.

If a swab is amylase positive, and PSA and KM negative, the decision on further testing depends on the location that the swab originated from (orifice versus non-orifice) and the sex of the victim. Refer to the Suspect Kit Processing Flow Charts for guidance.

If a swab is PSA and amylase negative, the decision on further testing depends on the location that the swab originated from (orifice versus non-orifice) and the sex of the victim. Refer to the Suspect Kit Processing Flow Charts for guidance.
Be sure to check for dried secretions with locations from the mouth, near the mouth, anal cavity, or near the anal cavity. Consult a supervisor to determine if a dried secretion from these locations should go on for extraction prior to kit closing.

**Vaginal and cervical swabs**

If a swab is sperm/PSA positive, make a second cutting from each positive swab for differential extraction.

If a swab is KM positive, consult with a supervisor

If a swab is sperm/PSA negative, consult with a supervisor.

**Oral and anal swabs**

If a swab is sperm/PSA positive, make a cutting from positive location for differential extraction.

If a swab is sperm/PSA negative, consult with a supervisor

18. After cutting all pertinent items, return all swabs and smears to their respective envelopes. Seal all kit envelopes with evidence tape and return to the kit. Seal the kit and return to a secure storage location.

**Suspect kit processing flow charts**
Suspect Kit – Oral, Anal, Vaginal, and Cervical Swabs

Is the victim female? 

Yes: Consult with a supervisor to determine if additional testing is needed

No: Sperm positive smear? 

Yes: Cut one sperm positive swab from each designated area for differential extraction

No: PSA positive? 

Yes: Return items to kit

No: Return items to kit – Serology report

Return items to kit
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Suspect Kit – Underwear (Suspect and/or Victim is Male)

- Does stain fluoresce or appear biological? No → Return items to lab – Semology report
  - Consult with supervisor to determine if additional testing is needed
- KM positive? No → AP positive?
  - PSA positive? Yes → Return items to lab
  - PSA negative? Yes → Amylase positive? Yes or No → Return items to lab
  - PSA negative? Yes → Amylase positive? Yes or No → Return items to lab
  - PSA negative? No → Amylase negative? No → Cut stain that is semen positive for differential extraction – autosomal
  - PSA negative? Yes → Amylase negative? No → Cut stain that is semen negative and either KM or amylase positive for non-differential extraction – autosomal
- AP positive? Yes → PSA positive? Yes → Return items to lab
  - PSA negative? Yes → Amylase positive? Yes or No → Return items to lab
  - PSA negative? No → Amylase negative? Yes or No → Return items to lab
- AP negative? No → Stain appears reddish/pink?
  - Yes → Cut stain that is semen negative and either KM or amylase positive for non-differential extraction – autosomal
  - No → Return items to lab – Semology report

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Suspect Kit – Underwear (Suspect and Victim are Female)

1. Stain appears reddish/brown?
   - Yes
     - KM positive?
       - Yes
         - Amylase positive?
           - Yes
             - Consult a supervisor and the case scenario to determine if additional testing is needed
           - No
             - Cut stain that is KM and/or amylase positive for non-differential extraction – autosomal
             - Return items to kit
       - No
         - Amylase positive?
           - Yes
             - Consult a supervisor and the case scenario to determine if additional testing is needed
           - No
             - Cut stain that is KM and/or amylase positive for non-differential extraction – autosomal
             - Return items to kit
   - No
     - Does stain fluoresce or appear biological?
       - Yes
         - Return items to kit – Serology report
       - No
         - Consult a supervisor and the case scenario to determine if additional testing is needed

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K. Evidence examination – non post-mortem exemplars

Follow the general guidelines for note taking and evidence examination when examining any exemplar item.

True exemplars:

An exemplar must have documentation stating that it is in fact from the person named. A “true exemplar,” such as a blood sample or an oral swab, will include paperwork from the MLI who obtained the sample, paperwork from the NYPD (including a voucher and sometimes a signed consent form), or paperwork from the DAO.

Use the General Packaging Worksheet for initial documentation of each item.

1. For a blood sample, follow the bloodstain preparation section of the Serology Manual. Cut a portion of the dried bloodstain card for exemplar extraction, using the initials of the individual in the short sample name.

2. For an oral swab, document the sample using the General Packaging and Swab Exam Worksheets. Cut approximately ¼ of the swab for exemplar extraction, using the initials of the individual in the short sample name.

3. Retain the victim exemplars from sexual assaults. Place the swab(s) in a coin envelope labeled with the FB number, voucher number, item number, victim name, analyst’s initials, and date of examination. The coin envelope should be placed in a Kapak envelope and heat sealed. The FB number should be written on the Kapak and the analyst’s initials and date of examination should be written across the seal. Place the exemplar in a secure storage location and return the empty packaging to the EU. For blood samples, retain the stain card and clean the empty tubes with 10% bleach and return them along with the packaging to the Evidence Unit.

Pseudo-exemplars:

It is the policy of the Department of Forensic Biology to accept and test “pseudo-exemplars”. It is our expectation that NYPD investigators will submit items with a reasonable probability of Controlled versions of Department of Forensic Biology Manuals only exist in the Forensic Biology Qualtrax software. All printed versions are non-controlled copies.
finding a single-source DNA profile from the suspect. The item must have been abandoned; common examples include a cigarette butt tossed in the street, a coffee cup left behind after questioning, or a bottle the suspect was seen handling. It is not acceptable to test items taken directly from a suspect (e.g. handcuffs for the DNA of the person that these were last used on) or items of evidence collected from an unrelated incident (e.g., bloody clothes from a suspect who was a victim of an assault).

1. Use the General Packaging Worksheet for initial documentation of each item.

2. For a cigarette butt “pseudo-exemplar,” document the sample using a Cigarette Butt Examination Worksheet. Cut a piece of the filter and paper portion for pseudo-exemplar extraction.

3. If an item (such as cup or bottle) is submitted, use the General Item Examination Worksheet for documentation. Use a cotton-tipped swab moistened with distilled water to swab the surface of contact. Briefly allow the swab to dry and then cut a portion of the swab for pseudo-exemplar extraction. Amylase testing is not necessary for pseudo-exemplars.

4. For other items submitted as pseudo-exemplars, cut or swab the item as appropriate. It may be necessary to consult with a supervisor to determine the best approach.

5. Remember to designate samples taken from pseudo-exemplars using an appropriate LIMS suffix to indicate that it is not a true exemplar. For example: “_AM” for bottles and cups or “_CB” for cigarette butts. For short sample description, include the item type and initials of the person providing the pseudo exemplar. For example: “bltRB” for a bottle or “cigRB” for a cigarette butt.

L. Evidence examination – condom

Condoms are often submitted to the Forensic Biology laboratory for examination. Follow the general guidelines for note taking and evidence examination when examining a condom.

Use an Evidence Packaging Worksheet for initial documentation of each item.
1. Describe the general condition of the condom (laid out flat, wadded up), color, and any trace evidence if present. If the condom was submitted “tied off,” document it as received then cut open for sampling.

2. If applicable, any stains must be documented by diagrams and/or photography. Note the location of the stain, size, heaviness (surface smear, etc.), and any directionality of the stain pattern. Each photograph must have a ruler visible in the frame, either a plain straight ruler or an x, y axis ruler.

3. Note whether fluids are present (liquid or dried). If the condom is found to be wet when opened, the item should be allowed to air dry after samples are taken. The item should not be heated or exposed to direct sunlight. If the item has become foul smelling, allow it to dry in the hood with the fan running.

4. Separately swab both the “inside” and “outside” of the condom, using no more than two swabs for each surface. Since it usually can’t be conclusively determined which surface is which, use quotes to describe the “inside” and “outside.”

5. Test both sets of swabs for the presence of blood, semen, and/or amylase as needed. Since the presence of a victim’s DNA on a condom can often be important, it may be necessary to perform DNA testing on a sample from a condom even if no blood, semen, or amylase is detected. Consult a supervisor if needed.

6. Do not sample a condom by cutting a portion of the condom.

M. Evidence Examination – Products of Conception

The term product of conception (POC) refers to either an embryo (up to the formation of organs in the first 8 weeks of gestation) or a fetus (up to approximately 30 millimeters and weighs approximately 4 grams).

The placenta is a temporary organ of pregnancy. Anatomically, the placenta has two parts: decidua (D), genetically identical to the mother, and chorionic villi (CV), genetically identical to the POC. Decidua appears as a compact tissue, while chorionic villi look more incoherent and loose. Morphological differentiation between D and CV can be made by observation:
By naked eye (Figure 1a and 1b)
• Using stereo-microscopy (Figure 2a and 2b),
• Using light microscopy of formalin fixed, paraffin embedded, and stained tissue (Figure 3a and 3b).

It is possible for tissues of POCs to lack uniformity, be of different gestational ages, or be differently preserved. Therefore, besides general guidelines for evidence examination, examination of POCs requires that some specific scenarios be taken into consideration.

Follow the general guidelines for note taking and evidence examination when examining POC.

Use a Product of Conception (POC) Packaging and Exam Worksheet for initial documentation of each POC item.

1. Describe the general condition of the item (full embryo/fetus, fragments, unrecognizable tissue parts, etc.).

2. Take one overview photograph of each item. Each photograph must have a ruler visible in the frame, either a plain straight ruler or an x, y axis ruler.

3. Weigh each item and document the tissue weight.

4. Determine if the POC is more or less than 24 weeks of gestational age (weight of ≥ 500g is considered > 24 weeks of gestational age).

5. Sampling of the item depends on the general condition of the item.
   a. If the POC is morphologically well defined, take a sample from it for DNA typing; the sample should be approximately 3x3x3 mm in size.
   b. If the POC is <24 weeks of gestational age and/or it is not morphologically well defined, rinse it several times in dH2O using Petri dish and observe it wet under MIDEO stereo microscope (following Protocol for Forensic Mitochondrial DNA Analysis, Section 4: MIDEO Macro/Microscopic Digital Imaging System, page 1-3).
Referring to Figure 2a and 2b for guidance, take a chorionic villi sample for DNA typing; the sample should be approximately 3x3x3 mm in size. If an exemplar from the mother/victim is not available, take a decidua sample as well.

c. If the POC is <24 weeks of gestational age, and/or it is not morphologically well defined, and/or morphological differences between maternal and fetal part of placental tissue could not be established using MIDEO as in step b above, take several samples from morphologically different regions and put them in separate embedding cassettes (Figure 4) for histological examination.

![Figure 4: Tissue Embedding Cassette](image)

Each sample should be approximately 10x10x5 mm in size. Close each cassette and label with a pencil. Submerge the cassettes in a prepared jar of formaldehyde. Cassettes, formaldehyde, and jars will be pre-provided by Histology Department.

After collection of all pieces is done, submit them to Department of Histology for further paraffin embedding, cutting, slide mounting and staining procedure. If needed, ask for consultation with a pathologist. Once the samples have been evaluated, follow the section of the Laser Microdissection procedure from Forensic Biology Protocol for STR Analysis (In Section 2: DNA Extraction). Make sure that the chain of custody is maintained.

d. If the POC is >24 weeks of gestational age, retain a sample for further testing. Inform OCME Identification Unit and keep the POC in a freezer, properly packed, until a permit for city burial is obtained by OCME Identification Unit. Return the empty packaging to the OCME Evidence Unit.
6. Submit samples for DNA extraction on an **Exemplar** test batch, using the notation “D” for decidual tissue and ACV® for chorionic villi as appropriate.

7. Depending on the outcome of the DNA testing, the disposition of the POC varies:

<table>
<thead>
<tr>
<th>Testing outcome</th>
<th>Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>No mother/victim exemplar, and DNA profile of the POC is female</td>
<td>- Retain the entire POC;</td>
</tr>
<tr>
<td></td>
<td>- Return the empty packaging to the OCME EU</td>
</tr>
<tr>
<td>No mother/victim exemplar, and DNA profile of the POC is male</td>
<td>- Retain a sample of POC for further testing;</td>
</tr>
<tr>
<td></td>
<td>- Dispose the remainder of POC in the red waste trash (If the POC is &gt;24 weeks old, follow step 5d);</td>
</tr>
<tr>
<td></td>
<td>- Return the empty packaging to the OCME EU</td>
</tr>
<tr>
<td>No mother/victim exemplar and DNA profile of the POC is a mixture</td>
<td>- Repeat testing (See Step 5 above)</td>
</tr>
<tr>
<td>There is a mother/victim exemplar and DNA profile of the POC is foreign to the</td>
<td>- Retain a sample of POC for further testing;</td>
</tr>
<tr>
<td>victim (mother), having expected allele sharing</td>
<td>- Dispose the remainder of POC in the red waste trash (If the POC is &gt;24 weeks old, follow step 5d);</td>
</tr>
<tr>
<td></td>
<td>- Return the empty packaging to the OCME EU</td>
</tr>
<tr>
<td>There is a mother/victim exemplar and DNA profile of the POC is a deducible</td>
<td>- Retain a sample of POC for further testing;</td>
</tr>
<tr>
<td>mixture</td>
<td>- Dispose the remainder of POC in the red waste trash (If the POC is &gt;24 weeks old, follow step 5d);</td>
</tr>
<tr>
<td></td>
<td>- Return the empty packaging to the OCME EU</td>
</tr>
<tr>
<td>There is a mother/victim exemplar and DNA profile of the POC is an undeducible</td>
<td>- Repeat testing, following Step 5a or 5b</td>
</tr>
<tr>
<td>mixture</td>
<td></td>
</tr>
</tbody>
</table>
For the return of empty packaging, bleach each container in which POC have been submitted using 10% bleach prior to return to the Evidence Unit.

Figure 1a: CV by naked eye

Figure 1b: CV by naked eye - detail
Figure 2a:
Stereo-microscopic (MIDEO) image of chorionic villi.

Figure 2b:
Stereo-microscopic (MIDEO) image of Decidua.

Figure 3a:
Microscopic image of formalin fixed, paraffin embedded and routinely stained decidua

Figure 3b:
Microscopic image of formalin fixed, paraffin embedded and routinely stained chorionic villi
N. Evidence Examination – Touched Items

Held or touched items may be expected to yield low amounts of DNA. These items should be swabbed or scraped according to the protocols described below.

1. Documentation

Record the Evidence Packaging as the initial documentation of each item.

a. Follow the evidence exam guidelines for proper documentation of all items and samples taken. For further clarification see below.

   i. Note the general appearance of the item. For example, note the color, the dimensions, and whether the item appeared to be dirty or possibly treated with latent print developers such as fingerprint powders or cyano-acrylate (fuming) etc.

   ii. Note the specific area being swabbed and/or any stains observed. Include the dimensions of the stain or area:

      a) If an area is reddish brown, KM test the area if appropriate. For a very small area, consult a supervisor.

b. Determine the areas of the item to be swabbed separately if necessary. Describe the sample assignment in detail in the notes. Examples follow:

   i. For duct tape used to bind a victim, multiple samples may be taken depending upon the circumstances of the case and the item. These samples may include the ends of the non-sticky side of the tape, the ends of the sticky side of the tape as well as the middle of the non-sticky side of the tape.

   ii. Similarly, a bat may be divided into the following three sections: the top or where the bat came into contact with the victim, the middle or barrel of the bat which may have the victim’s and/or the handler’s DNA, and the handle of the bat.

   iii. Each of the sections will be initially treated as separate samples.

2. Swabbing a touched item using SDS swabs

a. Obtain as many irradiated SDS swabs and aliquots of the 0.01% SDS swabbing solution as may be necessary for the item currently being examined. As a general rule, approximately 6 square inches may be effectively swabbed with one
SDS swab. This is dependent on the condition and type of evidence being examined.

b. Do not open the swab tube until you are ready to swab the item.

c. Clean a set of tweezers with 10% bleach, and 70% ETOH.

d. With a tube opener or lint-free wipe, open the tube and remove the swab with tweezers.

e. Dip a portion of the swab into the swabbing solution (0.01% SDS). Do not saturate, rather moisten, the swab. If too much SDS solution is used, DNA may be left behind on the item.

f. Swab the target area by folding or balling the swab up with the tweezers.

g. Thoroughly swab the target area with gentle pressure making sure to leave as little of the swabbing solution behind as possible.

**NOTE:** Multiple swabs may be used for a single area, as necessary. Document the use of multiple swabs and note the area which was swabbed. Only submit as many swabs in a single tube as may be effectively covered by digestion buffer (approximately 200µl) at the extraction stage. (The samples divided into separate extraction tubes may then be recombined into one extract in a microcon step.)

h. Should residual SDS be left on an item, use a dry SDS swab to collect it and include it in the extraction tube to be extracted along with the original swab(s).

i. Place the swab(s) into the extraction tube(s).

j. When swabbing more than one item from a case use a fresh tube of swabbing solution for each item.

k. Change gloves between items when swabbing different pieces of evidence.

3. **Cutting swabs submitted by another party**

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a. If evidence is a swab previously taken, cut the entire swab and place in an irradiated extraction tube.

b. Cut the exterior layer of cotton or surface of the swab that appears to have come in contact with the evidence. Make a cutting of one third of the swab as normal. Then, starting from the area of the initial cut, peel the outer layer of the swab. Cut in circular pattern, essentially lifting that top layer off the stick with the scissors. Take care not to cut the wooden stick.

Repackage the evidence and return to a secure storage location.

O. Evidence examination – Fingernail Scrapings (or Clippings)

Fingernail scrapings or clippings would be examined upon the request of the NYPD or law office and approval by a supervisor. Generally, information that indicates a struggle between the victim and the suspect must be provided in order to approve this testing.

Use the Evidence Packaging Worksheet for initial documentation, where applicable. In many cases, this may have been completed during the original examination of the sexual assault kit or post-mortem kit.

Note: Fingernail scrapings and clippings are to be sub-itemized by how they were received. Most often, they are initially separated by the right hand (containing scrapings or clippings or both) and the left hand (containing scrapings or clippings or both). For example, if the fingernails are item 1.4, they should be sub-itemized as items 1.4.1 (right hand) and 1.4.2 (left hand). Individual scraping dowels and fingernails must then be sub-itemized again before examination.

For example, if the fingernail packaging from a sexual assault kit contains all possible scrapings and clippings, the items should be listed as:

1.4 fingernail scrapings/clippings (itemized below)
   1.4.1 right hand fingernail scrapings/clippings (itemized below)
     1.4.1.1 right hand fingernail scrapings
     1.4.1.2 right hand fingernail clipping
     1.4.1.3 right hand fingernail clipping
     1.4.1.4 right hand fingernail clipping
     1.4.1.5 right hand fingernail clipping

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### Fingernail Scrapings

a. Complete a General Items Worksheet for the submitted fingernail scrapings. If packaged together, multiple scraping dowels can be examined at the same time (but sampled separately). If fingernail scrapings were received and previously documented in a sexual assault kit, you may need to edit the quantity and itemize the scraping dowels individually.

---

**Evidence Examination**

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<td>Quality Assurance Manager</td>
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</tr>
</tbody>
</table>

1.4.1.6 right hand fingernail clipping
1.4.2 left hand fingernail scrapings/clippings (itemized below)
   - 1.4.1.1 left hand fingernail scrapings
   - 1.4.1.2 left hand fingernail clipping
   - 1.4.1.3 left hand fingernail clipping
   - 1.4.1.4 left hand fingernail clipping
   - 1.4.1.5 left hand fingernail clipping
   - 1.4.1.6 left hand fingernail clipping

In cases where the right and left hands are packaged separately, one level of sub-itemization will suffice. For example:

2 right hand fingernail scrapings/clippings (itemized below)
   - 2.1 right hand fingernail scrapings
   - 2.2 right hand fingernail clipping
   - 2.3 right hand fingernail clipping
   - 2.4 right hand fingernail clipping
   - 2.5 right hand fingernail clipping
   - 2.6 right hand fingernail clipping

3 right hand fingernail scrapings/clippings (itemized below)
   - 3.1 right hand fingernail scrapings
   - 3.2 right hand fingernail clipping
   - 3.3 right hand fingernail clipping
   - 3.4 right hand fingernail clipping
   - 3.5 right hand fingernail clipping
   - 3.6 right hand fingernail clipping

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b. Cut a ~¼ inch piece from both ends of the individual dowel and place into one extraction tube (per dowel). Collect any debris that may have fallen off the dowel and place in an extraction tube.

c. Add the appropriate “_FN” suffix to all collected samples and submit for robotic extraction.

2. Fingernail clippings

a. Complete a Nail Examination Worksheet for each item. If packaged together, multiple fingernails can be examined at the same time (but sampled separately). If broken, pieces of fingernails should be treated as separate samples (there may be more than 10 samples).

b. Fingernails must be photographed since they will not be returned to their packaging. Fingernails can be grouped by hand for a photograph; photograph as described in the general guidelines of this manual.

c. Examine the fingernails under the stereoscope. Itemize any discovered skin or debris that can be separated from the fingernail as an additional sample.

d. KM test as needed. If a blood stain is suspected, collect the entire stain with a sterile swab moistened with water. Use a small piece of that swab for presumptive testing. If KM positive, consume the remainder of the collected sample for robotic extraction.

Note: With the exception of homicides, a KM positive sample is sufficient for the first round of testing. (For post-mortem samples, it is more likely that a KM positive is a result of the presence of victim, rather than foreign, blood.)

e. Cut longer fingernails in half; large samples may hinder the extraction process. Add the appropriate “_FN” suffix to all collected samples and submit for manual fingernail extraction.

Note: When submitting fingernails for extraction, the Evidence Item (fingernail) is “consumed”. For sexual assault kits, the empty packaging can be returned to the kit. For retained post-mortem samples, create a package in the LIMS and note it as “created in lab”. Add any remaining items to this package, print and affix the additional label. Post-mortem samples are to be retained in the appropriate post-mortem storage unit.
### EVIDENCE EXAMINATION

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</tbody>
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