CHAPTER 5 - ESTABLISHMENT INSPECTIONS

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There may be occasions where you may be accompanied on your inspection or investigation by other officials. These officials may be state or local officials who have their own inspectional authority or other officials who do not have authority to enter the firm. You should obtain permission from the firm’s most responsible person if officials without inspection authority wish to accompany you during your inspection/investigation. You should document in your EIR when other non-FDA officials accompany you during your inspection, and whether they entered under their own authority or the responsible individual at the firm gave permission (identify, by name and title, the responsible individual giving permission). See IOM 5.2.2 and 5.10.4.3.3.

5.1.1.1 - FDA Investigator’s Responsibility

Your authority to enter and inspect establishments is predicated upon specific obligations to the firm as described below. It is your responsibility to conduct all inspections at reasonable times and within reasonable limits and in a reasonable manner. Proceed with diplomacy, tact and persuasiveness.

During inspections or investigations, when you have evidence of conditions whereby there is a reasonable probability the associated products will cause imminent and serious adverse health consequences or death, you should notify your supervisor immediately to consider a Risk Control Review (RCR) evaluation.

5.1.1.2 - Credentials

Display your credentials to the top management official be it the owner, operator, or agent in charge. See IOM 5.2.2.

NOTE: Although management may examine your credentials and record the number and your name, do not permit your credentials to be photocopied. Federal Law (Title 18, U.S.C. 701) prohibits photographing, counterfeiting, or misuse of official credentials.

5.1.1.3 - Written Notice

After showing the firm’s representative your credentials, issue the original, properly executed, and signed FDA 482, Notice of Inspection, to the top management official. Keep the carbon copy for submission with your report. A notice of inspection is not required to be issued during foreign inspections; however credentials should be presented to the top management official.

5.1.1.4 - Written Observations

Upon completing the inspection and before leaving the premises, provide the highest management official available your inspectional findings on an FDA 483 - Inspectional Observations. See Section 704(b) of the FD&C Act [21 U.S.C. 374 (b)] and IOM 5.2.3 and 5.2.7.
5.1.1.5 - Receipts

Furnish the top management official the original of the FDA-484 - Receipt for Samples describing any samples obtained during the inspection. See IOM 5.2.4.

5.1.1.6 - Written Demand for Records

In low-acid canned food and acidified food EI's, an FDA 482a - Demand for Records (exhibit 5-2) is required under 21 CFR 108.35(h) and 21 CFR 108.25(g) to obtain records required by 21 CFR 113 and 114.

5.1.1.7 - Written Requests for Information

There are several methods of requesting records. These may include a request for information under LACF or AF inspections, 703 written requests, and requests for records under the BT Act (IOM 5.4.1.3).

5.1.1.7.1 – LACF / AF Food Inspections

In low-acid canned foods and acidified foods EI's, an FDA 482b, Request for Information (exhibit 5-3), is required under 21 CFR 108.35(c)(3)(ii) and 21 CFR 108.25(c)(3)(ii) to obtain information concerning processes and procedures required under 21 CFR 113 and 114.

5.1.1.7.2 – Requests for Records under Section 703 of the FD&C Act

Per CPG Sec. 160.300, Requests for Records under Section 703 [21 U.S.C. 373], evidence obtained in response to a specific written request under Section 703 cannot be used in a criminal prosecution of the person from whom obtained. With Supervisory approval, in certain circumstances, you may decide to issue a 703 written request when the importance of the evidence is crucial to protecting the public health.

Procedure: All 703 written requests must comply with IOM 4.4.7.2.2. Consider obtaining the evidence from other sources before using the 703 written request. In the case of foods and feeds, if there is a risk or threat of serious adverse health consequences, the district should invoke the BT Act records access authority. All BT Act records requests must comply with IOM 5.4.1.3.

5.1.1.8 - Business Premises

Authority to inspect firms operating at a business location is described in IOM 5.1.1 and requires issuing management an FDA 482, Notice of Inspection, and presenting your credentials. A warrant for inspection is not necessary unless a refusal or partial refusal is encountered or anticipated.

5.1.1.9 - Premises Used for Living Quarters

All inspections where the premises are also used for living quarters must be conducted with a warrant for inspection unless:

Owner Agreeable - The owner or operator is fully agreeable and offers no resistance or objection whatsoever or;

Physically Separated - The actual business operations to be inspected are physically separated from the living quarters by doors or other building construction. These would provide a distinct division of the premises into two physical areas, one for living quarters and the other for business operations, and you do not enter the living area.

In both the latter cases, proceed as any other inspection with the appropriate presentation of credentials and issuance of a Notice of Inspection.

5.1.1.10 - Facilities where Electronic Products are Used or Held

Section 537(a) of the FD&C Act provides the FDA with the authority to inspect the facilities of manufacturers in certain circumstances. The electronic product radiation control provisions were originally enacted as the Radiation Control for Health and Safety Act of 1968 (P.L. 90-602)

It is lawful for FDA personnel to enter the facilities of an electronic product distributor, dealer, assembler or user for the purpose of testing an electronic product for radiation safety when the entry is voluntarily permitted. Congress has not specifically prohibited FDA from conducting such voluntary examinations and such examinations would clearly agree with the congressional declaration of purpose expressed in section 532(a) of the RCH&S Act.

Under the Medical Device Authority, electronic products utilized in human and/or veterinary medicine, e.g., x-ray, laser, ultra-sound, diathermy, etc. can be considered prescription devices. In these cases the authority of Section 704 of the FD&C Act [21 U.S.C. 374] can be used to obtain entry to inspect the user facility. If the Medical Device Authority is utilized, credentials must be displayed and a FDA 482, Notice of Inspection, must be issued.

5.1.1.11 - Multiple Occupancy Inspections

You are required per FD&C Act 704(a)(1) [21 U.S.C. 374(a)(1)] to issue a Notice of Inspection, FDA 482, to each firm inspected. When firms have operations located in different sites or buildings, you should use judgment to determine when multiple FDA 482 forms need to be issued. For sites located a distance apart, it is preferable to issue a FDA 482 to the most responsible person at each site. One rule of thumb which can be used is if the sites or buildings are within walking distance, your original Notice of Inspection can be considered sufficient to cover both. During your initial interview with management, after you issue the FDA 482, make sure you clearly indicate the facility and sites you intend to inspect. The Act requires the issuance of a Notice of Inspection, but does not prohibit issuing multiple notices if management so requests. As
with all of our work, good judgment, and knowledge of the OEI and the FD&C Act are necessary in deciding what legally must be done.

5.1.1.12 - Authority for Examinations and Investigations

Section 702(a) of the FD&C Act [21 U.S.C. 372 (a)] authorizes examinations and investigations for the purpose of enforcing the Act.

5.1.1.13 - Authority to Implement Section 702(e)(5) of the FD&C Act

Section 702(e) of the FD&C Act [21 U.S.C. 372 (e)] contains certain authorities relating to counterfeit drugs including the authority to seize (“confiscate”) counterfeit drugs and containers, counterfeiting equipment, and all other items used or designed for use in making counterfeit drugs prior to the initiation of libel proceedings. This authority has been delegated, with certain restrictions, to holders of official credentials consistent with their authority to conduct enforcement activities. Additional authority in 702(e) to make arrests, to execute and serve arrest warrants, to carry firearms, or to execute seizure by process under Section 304 of the FD&C Act [21 U.S.C. 334] have not been delegated.

The agency does intend to utilize the authority contained in Section 702(e) to execute and serve search warrants, but such use does not require delegation from the ACRA.

Section 702(e)(5) contains authority for such delegated persons to confiscate all items which are, or which the investigator has reasonable grounds to believe are, subject to seizure under Section 304(a)(2). Items subject to seizure, and thus to confiscation under Section 702(e)(5), includes most things associated with counterfeit drugs. Confiscation authority does not, however, extend to vehicles, records, or items (i.e., the profits) obtained as a result of counterfeiting.

5.1.1.13.1 - SCOPE

Under this delegation, with supervisory concurrence and prior to the initiation of libel proceedings, investigators and inspectors are authorized to confiscate:

1. Any counterfeit drug,
2. Any container used to hold a counterfeit drug,
3. Any raw material used in making a counterfeit drug,
4. Any labeling used for counterfeit drug,
5. Any equipment used to make a counterfeit drug including punches, dies, plates, stones, tableting machines, etc.,
6. Any other thing which you have reasonable grounds to believe is designed or used in making a counterfeit drug.

NOTE: You and your supervisor must be constantly aware of the potential dangers involved in confiscating property from individuals. Special care should be taken to ensure your safety. Arranging for teams of investigators to conduct the investigation, or arranging for assistance by local police, or other agencies with police powers, should be considered in planning the confiscation of counterfeit materials.

5.1.1.13.2 - INSPECTIONAL GUIDANCE

Guidance provided for implementing the authority to confiscate drug counterfeits is as follows:

1. The authority is not to be utilized unless there has been an agency determination the drug to be confiscated is a counterfeit and it is a drug which "without authorization, bears a trademark, *** or any likeness" of a legitimate product. The determination usually is based upon evidence supplied by the firm whose product is being counterfeited. A written agency determination will issue to the District Director from the Office of Enforcement and Import Operations (OEIO), in conjunction with the Office of Medical Products and Tobacco Operations (OMPTO), and the Center for Drug Evaluation and Research.

2. When engaged in counterfeit investigations, you should proceed as follows upon encountering items to be confiscated.
   a. Evaluate safety needs and check the location to ensure it is safe to proceed. Do not attempt to remove an item by force. If it appears there will be resistance, contact the local police, or other agencies with police powers for backup, if not already done in advance.
   b. Inventory the items to be confiscated.
   c. Prepare a written receipt and offer it to the person in charge.
   d. Remove the items, if possible, from the premises (if they cannot be removed, secure them under seal).
   e. Place all items removed under lock at a secure location. In most cases, confiscated items will be stored at the district or resident post office until they are seized.

5.1.1.13.3 - FOLLOW UP GUIDANCE

After items are confiscated, certain actions must be taken to bring confiscated items under the control of the court.

Proceed as follows:

1. After an item is confiscated, immediately notify your supervisor.
2. Supervisors must then notify the appropriate compliance units of the items confiscated.
3. Compliance units should initiate seizure proceedings against any items confiscated.
4. OMPTO/Division of Medical Products and Tobacco Program Operations (DMPTPO) should be advised of any action utilizing this authority.

5.1.1.13.4 - SEARCH WARRANTS

Section 702(e)(2) contains authority to execute and serve search warrants.

Proceed as instructed by your district after a search warrant has been obtained.
5.1.1.14 - Products Imported Under the Provisions of Section 801(d)(3) of the FD&C Act

The FDA Export Reform and Enhancement Act of 1996 (PL 104-134 and 104-180) amended the FD&C Act by adding Section 801(d)(3) ("Import for Export") which permits the importation of unapproved drug and medical device components, food additives, color additives, and dietary supplements intended for further incorporation or processing into products destined for export from the United States. Section 801(d)(3) was subsequently amended by Section 322 of the Public Health Security and Bioterrorism Preparedness and Response Act of 2002 (Bioterrorism Act), Public Law 107-188, which specified certain requirements an importer has to satisfy in order to import a product under this Section. See IOM 6.2.3.4.

5.1.1.14.1 - REQUIREMENTS FOR BIOTERRORISM ACT

These requirements include:
1. A statement confirming the intent to further process such article or incorporate such article into a product to be exported,
2. The identification of all entities in the chain of possession of the imported article,
3. A certificate of analysis "as necessary to identify the article" (unless the article is a device), and
4. Executing a bond providing for liquidated damages in the event of default, in accordance with U.S. Customs.

This bond remains in effect until the final product is exported and destroyed.

In addition, the initial owner or consignee must keep records showing the use of the imported articles, and must be able to provide upon request a report showing the disposition or export of the imported articles. An article imported under this section, and not incorporated or further processed, must be destroyed or exported by the owner or consignee. Failure to keep records or to make them available to FDA, making false statements in such records, failure to export or destroy imported articles not further incorporated into finished products, and introduction of the imported article or final product into domestic commerce are Prohibited Acts under Section 301(w).

Filers making entry under the Import for Export provisions must either identify entry submissions with the OASIS Affirmation of Compliance "IFE" (Import for Export), or supply FDA with written documentation stating the product is entered under the Import for Export provisions. A Certificate of Analysis (as necessary) and identification of all involved entities must be submitted in writing to the import district. The import district will forward all written documents forwarded from the district where entry was filed. During the inspection examine the firm's records to determine the disposition of any items identified at time of entry as intended for incorporation into products for export. Document any instances in which such products were introduced into domestic commerce or cannot be accounted for (see IOM 6.2.3.4.3).

5.1.2 - INSPECTIONAL APPROACH

An establishment inspection is a careful, critical, official examination of a facility to determine its compliance with laws administered by FDA. Inspections may be used to obtain evidence to support legal action when violations are found, or they may be directed to obtaining specific information on new technologies, good commercial practices, or data for establishing food standards or other regulations. In order to facilitate on-the-job training, multiple points of view, and perspectives of firms being inspected whenever practical, those with assignment authority, should consider assigning different Investigator/s or different Lead Investigators at different times. This is recommended particularly when there have been multiple sequential NAI inspections or when the firm's management has been uncooperative.

The kind and type of inspection you conduct will normally be defined by the program, assignment, or your supervisor; according to the following definitions:

- Comprehensive Inspection - directs coverage to everything in the firm subject to FDA jurisdiction to determine the firms compliance status; or
- Directed Inspection - directs coverage to specific areas to the depth described in the program, assignment, or as instructed by your supervisor.

See IOM Subchapter 1.5 and 1.5.5 for information on safety, use of protective gear, trash disposal, dealing with potential hazards and other safety issues.

See special report requirements in IOM Subchapter 1.7.3 when objectionable conditions which may be of public health significance implicate establishments in other district(s).

5.1.2.1 - Depth of Inspection

The degree and depth of attention given various operations in a firm depends upon information desired, or upon the violations suspected or likely to be encountered. In determining the amount of attention to be given in specific cases, consider the:
1. Current Compliance Program,
2. Nature of the assignment,
3. General knowledge of the industry and its problems,
4. Firm history, and
5. Conditions found as the inspection progresses.

5.1.2.2 - Inspection Walk Through

A walk through inspection of the premises should be conducted as early as possible to become familiar with the operation and to plan the inspection strategy. A walk through visual inspection of the manufacturing site is helpful in establishing the depth of the inspection, learning about products and processes, identifying sources of manufacturing records and identifying potential areas of concern. The size of the facility, the number of employees, employee practices, environmental conditions inside and outside the plant, raw materials, manual and automated processes, sources of contamination, manufacturing flow, method of data collection including computer terminals, are some of the areas to be taken into consideration in establishing the depth of the inspection. A visual inspection of a manufacturing site should also be used to check obvious potential problem areas such as: general housekeeping, state of operation for processes and processing equipment, and people dependent operations. Visual inspections of areas used for failure investigation, product sampling and testing, product reworks, return goods, and product quarantine areas should be inspected for obvious potential product problems.

Depending on the product being inspected, some of the general inspectional equipment an investigator should have available, may include, eye and ear protection, boots and protective clothing. Some specialized equipment may include radiation or EO monitoring devices, magnifiers, and timing devices as needed. For some domestic and foreign plant sites, investigators may be required to be inoculated prior to the inspection for protection from potential environmental concerns such as hepatitis, yellow fever, malaria and live biological products which may be encountered in vaccine products. See subchapter IOM 1.5.

5.1.2.3 - Signing Non-FDA Documents

Occasionally a firm will request you sign various documents including:
1. A waiver which will exempt the firm from any responsibility or liability should an accident occur and you are injured on the firm's premises,
2. Form letters concerning access to confidential information the firm does not want released,
3. Information/data you request during the inspection be put into writing, etc.

If you receive such a request, inform the firm you are not authorized to sign such documents, letters, requests, waivers, etc., but will report the firm's request in your EIR. The use of common sense is expected with this procedure. All FDA employees are authorized to sign-in and sign-out at a firm and to comply with security measures employed by the firm, including documenting the removal/replacement of seals to inspect vehicles and containers. See IOM 4.3.4.3 and 4.5.4.6. Obviously, the key issue is you are not authorized to waive, without supervisory approval, any of FDA's rights to inspect, sample, photograph, copy, etc. or to sign any interstate shipping record document which could infer the firm could not be prosecuted under the Act.

5.1.2.4 - Technical Assistance

If you determine specialized technical assistance is necessary in conducting inspections of new technologies, products or manufacturing procedures, it may be available through Regional or National experts, other ORA components or Center scientists and engineers. If specialized skills are necessary and are not available locally or through your Region, contact the Division of Food and Feed Program Operations and Inspections (DFFPOI) for CFSAN and CVM products or the Division of Medical Products and Tobacco Program Operations (DMPTPO) for CBER, CDER, CDRH and CTP products. See FMD 142 and IOM 1.9.2.2.1 for additional information.

5.1.2.5 - Team Inspections

The use of teams to conduct inspections may be beneficial. Very often individuals well versed in an analytical or inspectional technique or technology can provide assistance and advice.

When inspection teams are involved in an inspection, one investigator will be designated as the team leader by the inspecting district or by DFFPOI or DMPTPO if a headquarters directed special inspection is involved. The team leader is in charge of the inspection and bears the overall responsibility for the inspection and the EIR. A team may consist of multiple investigators, laboratory personnel and other FDA employees, and your supervisor/coach, who may participate as part of the ORA Quality Assurance program.

5.1.2.6 - Post-Inspectional Contacts

If the firm contacts the Investigator after the inspection regarding the inspection or follow-up, the Investigator should refer the request to his or her supervisor or to Compliance Branch if a regulatory action is contemplated. The Investigator should not respond to the firm regarding the adequacy of the firm’s response to inspectional observations or any follow-up planned.

After the inspection is concluded, if the Investigator finds that a document or other required information is missing, the Investigator should discuss the needed information and how to proceed with their supervisor.

5.1.2.5.1 - TEAM MEMBER RESPONSIBILITIES
Each team member is responsible for preparing those portions of the report pertaining to his/her activities. Team members shall identify their portion of the report so they can later identify that portion as the part he/she performed and reported. Since reports should be written in the first person, one system might be to head each portion with a statement "The following operation(s) was/were observed and reported by Investigator ____________", who can then report in the first person.

All team members must sign the original EIR. Ideally, all team members should sign the FDA 483, if one is issued. However, issuance of the FDA 483 should not be delayed, in the absence of a team member's signature. See IOM 5.2.3 for instructions for signing a multi-page FDA 483.

5.1.2.5.2 - TEAM LEADER RESPONSIBILITIES

The Team Leader shall be responsible for:

1. Issuing unused notebooks for taking regulatory notes during the E1 or investigation to headquarters personnel on the team. He/she is also responsible for instructions on their use, if necessary, and when the report is finished, for obtaining the headquarters individual's signature on the original EIR and completed and properly identified regulatory notes and submitting them to the supervisor for filing. See IOM 2.1.3.

2. Directing the overall inspection to accomplish the objectives of the assignment including;
   a. Planning the inspection,
   b. Scheduling and coordinating team members' pre-inspection preparations,
   c. Determining, to the extent possible, the firm will be open and operating,
   d. Planning for needs of visiting scientists if applicable. When the team leader is not familiar with all the processes or technology involved in the inspection, provide for primary coverage of selected areas by other team members,
   e. Determining an orderly, efficient, and effective approach and sequence to be used and discussing the inspection plan with the team,
   f. Modifying the inspection plan as necessary during the E1, to permit following leads, documenting evidence, etc.,
   g. Setting team policy on how communications with the firm are to be handled,
   h. Discussing personal conduct in dealing with headquarters personnel as necessary,
   i. Assuring an early understanding by team members of their roles in note taking and reporting,
   j. Assuring communications are open among team members, especially if the team is allowed to separate and work independently,
   k. Reviewing inspection progress at least daily, discussing remaining objectives with the team members, and setting objectives for the following day,
   l. Continuously assessing the progress of the inspection to evaluate how the inspectional approach is working and to keep the district supervisor advised of the inspection's progress,
   m. Providing guidance and direction to team members as necessary,
   n. Advising each team member of reporting responsibilities and dates when drafts are to be provided,
   o. Following up promptly on any delays or failures to report as required, and
   p. Assisting the supervisor with further follow up, as indicated.

3. Making sure any person who joins the team after the inspection has started presents credentials and issues a FDA 482, Notice of Inspection to the firm prior to actually taking part in the EI;

4. Completing and/or correcting the computer generated coversheet;

5. Preparing the Summary of Findings;

6. Completing all headings of an administrative nature in the narrative report;

7. Compiling and submitting the complete final report; and

8. Resolving any disputes or differences of opinion among the team members, including items, which may be listed on the FDA 483.

5.1.3 - INSPECTION OF FOREIGN FIRMS

Inspectional requirements apply to all inspections, including foreign inspections. However, there are some exceptions. For instance the FDA 482 is not issued, unless the firm is a U.S. Military facility. Be guided by relevant Compliance Programs and the Guide to International Inspections and HHS Travel Manual for other differences.

5.1.4 - INSPECTIONAL PRECAUTIONS

Our concern over microbiological contamination emphasizes the need for you to be alert to criticism or allegations that you may have contributed to or caused contamination at a firm. This is especially important in drug firms and high-risk food firms, among others. You must adhere to good sanitation practices to refute any such criticisms. You could also unknowingly introduce or spread disease during inspections of or visits to animal production or sale facilities, conducting environmental investigations at poultry layer facilities, conducting dairy farm inspections or audits of state activities, investigating tissue residue reports or working in the veterinary bioresearch area. See IOM 5.2.10 for information outlining precautions for you to follow.

Exercise caution in all activities in the firm. Follow the firm's sanitation program for employees and wash and sanitize hands, shoes, vehicles and equipment as indicated. Restrict unnecessary movement between various areas in plants and when possible, complete your activities in one area before moving to the next.

When inspecting areas where sterility is maintained or sterile rooms are located (especially in pharmaceutical or device firms), follow the sterile program required of the firm's employees. In general it is unnecessary to enter sterile rooms except in the most extraordinary circumstances. These areas are usually constructed to provide
visual monitoring. Take no unsterile items with you (notebook, pencils, etc.). In this type of situation you can enter your observations in your regulatory notes immediately after leaving the sterile area.

Always use aseptic techniques, including hand sanitizing, when collecting in-line and raw material samples, as well as finished product samples for microbiological examination. See IOM 4.3.6.

Do not use or consume a firm's products at any of a firm's facilities. This could be interpreted as accepting a product as being satisfactory and could possibly embarrass you and the Agency, both during the inspection and in the future. In general, consuming food products in a manufacturing area is considered an objectionable practice.

When conducting inspections of firm's using chemicals, pesticides, etc., ask to review the Material Safety Data Sheets (MSDS) for the products involved to determine what, if any, safety precautions you must take. This could include the use of respirators or other safety equipment.

5.1.4.1 - Clothing

Wear clean coveralls or other protective clothing for each inspection and if circumstances dictate, use a clean pair when returning from lunch, or upon entering certain machinery or critical areas.

Remove and secure all jewelry, pens, pencils, notebook, etc., so they cannot fall into the product or machinery. Do not depend on clips on pens, etc., to hold these items in your outer pockets.

Clean protective clothing should be either individually wrapped or placed in clean plastic bags and taped to protect from contamination. If the package has been sterilized, protect the package from possible contamination or puncture. The package should not be opened until you are ready to use the clothing. After use, clothing should be turned inside out as it is removed, and immediately placed in clean paper or plastic bags to prevent spread of contamination until washed and/or sterilized.

Use disposable hair and head coverings throughout the inspection and disposable hand and foot coverings in areas where floor tracking or cross contamination may be a factor. Use hard hats and other protective devices where the situation dictates.

If reusable protective boots are used, wash and sanitize before each use. Always use sterile disposable boot covers when entering machinery such as dryers or where unavoidable contact with product is a factor.

When discarding contaminated disposable head and boot coverings, it is suggested they be placed with used clothing for proper disposal after leaving the plant area.

See IOM 5.2.10.1 for protective clothing and equipment necessary when visiting livestock or poultry producing areas.

5.1.4.2 - PHS Recommendations - Basic Sanitary Practices

FDA personnel are not required by law to have health certificates, take physical exams or submit to requirements, which ensures their compliance with sanitary procedures in the performance of their official duties. However, it is critical you adhere to basic sanitation practices. See IOM 1.5.1.5.

The Food Code 2013 is available electronically from the FDA CFSAN web page under Federal/State Programs-Retail Food Safety References. Printed copies may be ordered from the National Technical Information Service website.

5.1.4.3 - Representatives Invited by the Firm to View the Inspection

While conducting an inspection, you may find the firm's management has invited individuals who are not directly employed by the firm to view the inspectional process (e.g., representatives from the press, trade associations, consumer groups, congressional staff, other company officials).

Regardless of whom the firm invites to observe the progress of an inspection, the presence of outside representatives should not disrupt the inspectional process. You should continue to conduct the inspection in a reasonable fashion. The presence of these individuals should have no impact on the manner in which the inspection progresses except you should take precautions to preserve the confidentiality of any information you may have obtained as a result of the Agency's statutory authority. This is especially true when the inspection is recorded via videotaping, other photography, and/or audio recordings. Where applicable, refer to IOM 5.3.5 for procedures on how to prepare your own recording in parallel with the firm's recording.

It is the Agency's position that while the investigator must protect privileged information provided to him/her during the inspection, it is the firm's responsibility to protect privileged/confidential information observed or recorded by those individuals invited by the firm.

5.1.5 - GENERAL PROCEDURES & TECHNIQUES

The procedures and techniques applicable to specific inspections and investigations for foods, drugs, devices, tobacco products, cosmetics, radiological health, or other FDA operations are found in part in the IOM (inspectional and investigational policy/procedure), various Guides to Inspections of... (a "how to" guidance series), and the Compliance Program Guidance Manual (program specific instructions). Some procedures and techniques which may be applicable to overlapping areas or operations are as follows:

5.1.5.1 - Candling
Candling is defined as: "to examine by holding between the eye and a light, especially to test eggs in this way for staleness, blood clots, fertility and growth." Like most techniques learned through the food inspection programs, there are uses for this technique in other program areas such as looking for mold in bottled liquids which could be drugs, devices or biologics. Candling can also be useful in the examination of original documents to see below-whiteout or to look for over-writing.

Many types of products lend themselves to inspection by some type of candling. For these products, firms generally have candling equipment which may be built into the production lines or may be a separate operation.

Where checking products by candling, it may be possible to utilize the firm's candling equipment. Various other light sources for candling are also available including overhead projectors. Exercise care when using overhead projectors and protect the glass surface and the lens from scratches and damage. All candling is best accomplished when light outside the item being candled is masked so the light passes through the object rather than being diffused around it. A heavy paper or cardboard template can be quickly prepared at the time candling is done.

5.1.5.2 - Label Review

Do not undertake a critical review of labels unless instructed by the assignment, program, or your supervisor. Limit your comments to the mandatory label requirements required by the Acts. However, if after review of the formula, it is obvious an active ingredient or an otherwise mandatory ingredient statement does not appear on the label, such discrepancy may be called to management's attention. See also IOM 5.2.3.2 regarding labeling for blood and blood products.

If asked for other label comments, refer the firm to the appropriate Center to obtain a label review.

When the labeling is suspect or when you are requested to collect labels/labeling, collect three copies of all labels and accompanying literature for further review. For medical devices, if there is a question regarding the need for a new 510(k) or PMA supplement, it is essential the label and labeling be collected.

5.1.5.3 - Field Exams

A field examination is an on-site examination of a domestic product (or a foreign product in domestic channels of trade) sufficient in itself to determine if the product is in compliance with the Acts enforced by FDA. A field exam can be conducted of any commodity in any location. If the examination does not reveal a violation or the appearance of a violation, a sample of the lot is usually not collected. If your exam reveals a violation or potential violation, you should collect an official sample. With the implementation of FACTS, your time spent conducting the field exam is reported even if you do collect a sample. Only the actual time spent in the collection of the sample would be reported as sample time.

Instructions on how to conduct a field exam are contained in "Guides to Inspection of *** and Compliance Programs. The Sample Schedules in Chapter 4 also provide guidance on lot examinations for special situations.

SUBCHAPTER 5.2 - INSPECTION PROCEDURES

5.2.1 - PRE-INSPECTIONAL ACTIVITIES

Prior to the start of any inspection or investigation, you should conduct a number of activities. These will differ based on whether this is an inspection or an investigation. You should review the establishment's factory jacket (if one exists), and registration and listing (if applicable) information. The purpose of this review is to determine the location of the establishment and obtain an overview of the establishment's operations and products as well as an understanding of their compliance history. You should also review the establishment factory jacket to determine if there were any prior safety issues noted, e.g. documented Investigator safety incidents or whether any specific personal protective equipment is needed prior to the start of the inspection. If there has been a past personal safety incident, you should discuss with your supervisor and develop a Situational Plan prior to the start of the inspection. See IOM 5.2.1.4 – Personal Safety Plan.

Prior to initiating any inspection you should become familiar with the reporting requirements for the specific assignment, as well as the requirements of IOM Subchapter 5.10.

If the inspection or investigation is a directed assignment from a Center, ORA headquarters or another district, read the assignment and attached materials to assure you understand the assignment. If the inspection or investigation is being conducted in part or solely as a recall follow-up or complaint, refer to Chapter 7 (Recalls) or Chapter 8 (Investigations) of the IOM for additional guidance.

You should review the applicable FACTS assignment to determine if the Personal Safety Alert indicator is checked for this specific firm. The reason for the Personal Safety Alert should be listed in the Endorsement and should be accompanied by a Memo to the Establishment File Jacket or documented in a prior EIR. See IOM 5.2.1.3 FACTS Personal Safety Alert.

You should also review the applicable Compliance Program Guidance Manual(s) prior to the start of your inspection or investigation. ORA's Division of Medical Products and Tobacco Program Operations (DMPTPO) and Division of Food and Feed Program Operations and Inspections (DFFPOI) have written numerous Inspection Guides to assist you in conducting inspections of various types of establishments, products or processes. You
should become familiar with the appropriate guides prior to the start of the inspection and utilize them as needed throughout the inspection. The Centers have issued numerous guidance documents for industry. These documents are normally posted to the appropriate Center's Internet web site.

Subchapters 5.4-5.9 of the IOM contain additional, program specific pre-inspectional activities, which you should follow.

Imported products cross all program areas and our regulation of them does not stop at the border. Determine if there are any "import for export" follow-up assignments and be prepared to cover them during your inspection. See IOM 6.2.3.4 for guidance. Please be alert to imported products whenever you make an inspection. During inspections of domestic firms, if you encounter imported products that appear adulterated, misbranded, counterfeit, tampered with or otherwise suspect, attempt to fully identify the product and the source of the imported products. Contact your supervisor and Division of Import Operations (DIO) if necessary.

5.2.1.1 - Pre-Announcements

Pre-announcements are mandatory for all medical device inspections in accordance with the criteria and instructions below and some BIMO inspections. In all other program areas, pre-announcements may be made at the discretion of the district. If you are going to visit facilities where livestock (including poultry) or wild animals are housed or processed, review IOM 5.2.10. In general, it may be inappropriate to pre-announce inspections of food establishments, blood banks, source plasma establishments and some BIMO inspections, but this too is subject to district discretion. If a district believes pre-announcing an inspection of an establishment will facilitate the inspection process then the procedures below for doing pre-announcements for medical device inspections should be followed. ORA's primary purpose for pre-announcing is to assure the appropriate records and personnel will be available during the inspection. It is not to make an appointment for the inspection. It should not be referred to as an appointment to inspect. When doing a pre-announcement, it is important you communicate to the establishment the purpose of the inspection and a general idea of the records you may wish to review. If you find neither the appropriate personnel nor records available, note this in your Establishment Inspection Report (EIR).

In the case of drug inspections, if efforts to schedule a pre-announced inspection are met with unreasonable delays by the establishment, including requesting a later start date without a reasonable explanation, it may constitute a delay of an inspection under section 501(j) of the FD&C Act [21 U.S.C. 351(j)]. FDA will make reasonable accommodations for local conditions such as weather, holidays, or, where appropriate, manufacturing campaign schedules. However, if faced with an unreasonable delay by the establishment, you may call the responsible person's attention to 501(j) of the Act. Talk with your supervisor to determine whether the length of a particular delay may be considered unreasonable, even in cases in which the explanation given for the delay may be reasonable.

The District may use this data in the future when considering whether this establishment should be eligible for pre-announced inspections.

The following is the general outline for pre-announcement of medical device inspections. You are advising the establishment's management of the date and time you will be arriving at the establishment to conduct the inspection. The establishment has no authority to negotiate this. If you, as the investigator, feel the need to accommodate the establishment's request, be sure there are sound reasons for doing so and report them in your inspection report.

5.2.1.1.1 - BASIC PREMISES

Pre-announcement of inspections is to be applied only to establishments that meet specific criteria. Pre-announcement may be considered for establishments that manufacture both drugs and devices or biologics and devices. The eligibility of an individual establishment for pre-announced inspection is at the discretion of the inspecting office using clearly described criteria. (See Criteria for Consideration) The district does not have the discretion to decide the types of medical device establishments eligible for pre-announcement, but may decide the specific establishments' eligibility because they meet the criteria.

The pre-announcement should generally be no less than 5 calendar days in advance of the inspection. Should a postponement be necessary, the decision as to rescheduling rests with the investigator/team, but the new inspection date should not be later than 5 calendar days from the original date. Inspections may be conducted sooner than 5 calendar days if requested by or acceptable to the establishment and if this date is acceptable to the investigator/team.

To participate in the pre-announcement portion of the program, establishments are expected to meet the commitment to have appropriate records and personnel available during the inspection.

Pre-announced inspections will not limit an investigator's authority to conduct the inspection. Inspections will be as thorough as necessary.

5.2.1.1.2 - CRITERIA FOR CONSIDERATION

When deciding whether an establishment qualifies for a pre-announced inspection, you must consider whether both the type of inspection and the establishment's status meet the following specific criteria.

5.2.1.1.2.1 - Type of Inspection

Only the following types of inspections are appropriate:
1. Pre-market inspections (PMA, 510(k))
2. Foreign inspections
3. Quality System/Good Manufacturing Practice (QS/GMP) inspections:
   a. Biennial routine inspections
   b. Initial inspections of new facilities or newly registered establishments
   c. Initial inspections under new management and/or ownership.

5.2.1.1.2.2 - Eligibility Criteria

Establishment's eligible for pre-notification should meet the following requirements:
1. Non-violative QS/GMP inspection histories (inspections classified as no action indicated (NAI) or voluntary action indicated (VAI)). For VAI, adequate corrections of conditions observed and listed on FDA 483 during the previous inspection were verified and did not lead to any further agency action.
2. To remain eligible for pre-announced inspections, establishments must have a history of having individuals and/or documents identified in previous pre-announced inspections reasonably available at the time of the inspection.

5.2.1.1.3 - PROCEDURES

Procedures:
1. The investigator designated to conduct the inspection will contact the most responsible individual at the facility. You should leave a message requesting a return call if the most responsible person at the facility is unavailable at the time the call is made. The district should use good judgment as to what is a reasonable time frame to await the return call.
2. Changes in dates should be kept to a minimum. If a change is made, a new date should be provided as soon as possible, which will facilitate the inspection and accommodate the investigator’s schedule. The establishment should provide a valid reason for requesting a change in the start date. A valid reason should be the same as you would accept if presented with the information during an unannounced inspection.
3. Inform the establishment as to the purpose, estimated duration, and the number of agency personnel expected to take part in the inspection. The products or processes to be covered should be described if this will facilitate and be consistent with the objectives of the inspection.
4. When known, specific records/personnel will be requested at the time the inspection is pre-announced.
5. The notification should be as specific as reasonably possible and specify the date for the start of the inspection.

Include in your EIR whether or not the inspection was pre-announced and include information on any difficulties experienced in notification or accessing records or personnel, which should have been available as a result of pre-announcing the inspection. For medical device establishment inspections, if not pre-announced, describe briefly in the EIR why not. If an establishment should become ineligible for pre-announcement, the endorsement of the EIR should include this statement. This information will be necessary for making a determination regarding future pre-announced inspections of the establishment. In addition, it is advisable to inform the establishment during the current and subsequent inspections of the action(s), which may have caused them to be ineligible for pre-announcement.

Subchapters 5.4-5.9 of the IOM contain additional, program specific pre-inspectional activities, which you should follow.

5.2.1.2 - Personal Safety

ORA considers the safety of investigators, inspectors and all those who meet with regulated industry to be of the utmost importance. Personal safety concerns are defined as those factors FDA employees should maintain awareness of which potentially affect their safety during an inspection, such a threatening situation; or where specific personal protective safety equipment is warranted; or where a particular inspection may be medically contraindicated for specific FDA personnel. When these conditions are noted during an inspection, the investigator should discuss the situation with their supervisor and ensure that the Personal Safety Alert is checked in FACTS and a Memo to the File is generated – see IOM 5.2.1.3. For information concerning personal protective equipment, see IOM Subchapter 1.5.

Physical resistance to FDA inspections and threats to, or assaults on, FDA employees engaged in their work are extremely rare. However, there will be times you are confronted by unfriendly or hostile persons. ORA has offered various conflict resolution training courses to assist and prepare you for how to diffuse a situation. In most instances, conducting your activities with tact, honesty, diplomacy, and persuasiveness will be enough to diffuse the situation. While at times, you may have to adopt a firm posture, you should not resort to threats, intimidation, or strong-arm tactics. Refer to IOM 5.2.5.4 for Hostile and Uncooperative Interviewees.

Safety is the responsibility of all FDA employees, including you, your supervisor and other Agency management. When you receive an assignment, it is important to evaluate the assignment not only in accordance with IOM Section 5.2.1, but also with respect to your personal safety. If you determine there is the possibility of a threat to your personal safety, consult with your supervisor. You and your supervisor should consider developing a Situational Plan in preparation for the inspection.

5.2.1.2.1 - PREPARATION

Below are some suggested items the District may consider when preparing for your next assignment to assess if
there are potential personal safety issues. This list is not meant to be all inclusive.

1. Does the assignment involve working with other Federal Agencies such as U.S. Marshals, Federal Bureau of Investigations, and U.S. Customs in executing search warrants, seizures, etc.?

2. Does the assignment involve working with or contacting FDA’s Office of Criminal Investigations (OCI)?

3. Does the assignment involve a firm where there is a suspicion and/or knowledge of questionable or illegal activities?

4. Does the assignment involve a suspected tampering and/or a visit to an individual’s residence?

5. What is the past history from a personal safety standpoint with the prior interactions with representatives of this firm? Have the FDA’s state counterparts or other Federal and/or local agencies indicated a concern for personal safety? What does the firm’s establishment file indicate about personal safety over the past inspections?

6. What is the location of the firm or the operation? Is it in an area which may be unsafe? Have the inspected firm or any of its employees been uncooperative with government officials?

7. Is the firm known to the Agency? Has the Agency any additional information which would assist in your evaluation?

If these questions and/or others result in a concern for your personal safety, then a Personal Safety Plan should be developed and approved by district management before conducting the assignment. See IOM 5.2.1.4 – Personal Safety Plan.

Due to the unlimited variability of potential safety situations, it is not feasible to prescribe in the IOM what to do in every instance. The decision of what to do in each individual circumstance rests with the investigator and their district management. Your district management is most familiar with the specific firm in question, the regulated industry, as well as other local Federal, State and Local officials who may be able to provide you additional information and assistance. In addition, the experience of your district management combined with the various training courses on conflict resolution may also be consulted. Districts should notify the Office of Operations (OO) (Gail Katz) to inform headquarters of any potential safety concern, so that personal safety issues may be tracked. OO will also maintain a library of Personal Safety Plans which may also be of use to your District. OO may be contacted at 301-796-0358 or at the following personal safety e-mail address: ORAHQOMPTOCOSAF@FDA.HHS.GOV.

5.2.1.2.2 - PHYSICAL RESISTANCE/THREATS/ASSAULTS

If you receive physical resistance or threats, or if you sense the real possibility of an assault, disengage from the confrontation, get to safety, and call your supervisor immediately. Make careful and exact notes later of who said what to whom, who did what, and whether someone tried or succeeded in threatening, assaulting or taking information or equipment or samples from you. Be careful in any descriptions you give or write of such events, just as you are in recording other evidence that may result in a court case. Your safety is more important to the United States than the inspection or the sample. FDA will work with law enforcement government officials, e.g., the Federal Protective Service (FPS), FDA’s Office of Criminal Investigations’ (OCI) Special Agents, local police, or United States Marshals to assist an inspection team if there is a reasonable fear of danger to the investigator.

If you are assaulted (either physically or put in fear by threats of physical violence), your supervisor can summon local police, the Federal Protective Service (1-877-437-7411), United States Marshals, FBI or contact OCI headquarters for assistance (301-294-4030). While OCI does not normally provide physical security in these cases, they will assist in threat evaluation based on specific facts and available criminal databases. OCI can also make contacts with local police and federal agencies based on previous established liaisons. If you have been assaulted or threatened and you are unable to reach your supervisor or other district management, you should contact the local police in the area where the assault or threat occurred. Be careful in any descriptions you give or write of such events, just as you are in recording other evidence that may result in a court case. Make sure that any inspected facility where weapons are observed, or where threats or assaults occur, is identified on that facility’s Endorsement page of the inspection report for that facility and to your supervisor, so that Investigators or Agents who follow you into that facility will be alert to those possibilities. Your supervisor would also be responsible for checking the Personal Safety Alert box in FACTS and for beginning the notification process to alert other Federal or State agencies that also inspect the facility of the possible danger. For more information see IOM 5.2.1.3 Personal Safety Alert. For specific safety guidance related to inspections and interviews, see IOM 5.2.5.4.2 Hostile and Uncooperative Interviewees.

In addition, in any instance where you have perceived a threat to your personal safety during an inspection, investigation or sample collection, you should exit the situation immediately and report it to your supervisor. You should then write a memorandum of the event in a factual manner including information pertaining to the who, what, when, where, and how of the event. Be careful in any descriptions you give or write of such events, just as you are in recording other evidence that may result in a court case. This memo will be filed in the official establishment file jacket and copies be sent to any and all resident posts and import offices who may interact with this firm. The memo will be filed on the opposite side of the folder from all other documents and will be a printed on eye-catching color paper in order for the document to be visible to the next Investigator. The memo should be retained and maintained within the district. A copy of the Memo documenting the personal safety situation should also be sent to Office of Operations, Attn: Gail Katz.
5.2.1.3 – FACTS Personal Safety Alert

Within the Maintain Firms Option in the FACTS system, there is Personal Safety Alert option that allows the supervisor (FACTS Supervisor Role) to check the appropriate box to advise the FDA investigator that there is a personal safety issue. Only the FACTS Supervisor Role will allow for updating the Maintain Firms screen. This personal safety alert may be selected when there is a potential hazard identified:

1. Where a previous threat/assault or physical resistance occurred
2. Where specific personal protective equipment is needed (respirators, etc.)
3. Where there are specific medical considerations for a population of investigators (e.g. the firm manufactures a drug hazardous to women of child-bearing years or those with allergies to peanuts, penicillin, or other products.)

In any example listed where there is a Personal Safety Alert, the specific safety alert should be documented both in the Endorsement and in a Memo to the File. The memo should be flagged “MEMO TO FILE - PERSONAL SAFETY ALERT” and should provide the factual information to support why the investigator should be alerted to the safety issue. Be careful in any descriptions you give or write of such events, just as you are in recording other factual evidence that may result in a court case. The memo should be filed in the official establishment file jacket and copies sent to any and all Resident Posts and Import Offices who may interact with the firm. The memo will be filed on the opposite side of the folder from all other documents and will be a printed on eye-catching color paper in order for the document to be visible to the next Investigator. The memo should be retained and maintained at the District Office. A copy of the Memo documenting the personal safety situation should also be sent to the Office of Operations, Attn: Gail Katz. The supervisor and/or other district management will be responsible for evaluating any corrective actions taken by the firm or individual to remove or stop the potentially dangerous situation or condition. If the situation remains potentially dangerous, the Personal Safety Alert should be maintained in FACTS. Follow-up inspections at the facility should continue to document whether or not the safety situation continues exists. If the situation has been resolved (new management, dismissal of an employee, cessation of penicillin in a facility, etc.) the Personal Safety Alert should be removed from FACTS by the supervisor.

5.2.1.4 – Personal Safety Plan

A Personal Safety Plan is an investigative tool developed to assist in managing and preparing for a potentially dangerous situation. Districts should consider developing a Personal Safety Plan when the conditions surrounding the specific inspection, investigation or sample collection indicate a plan is needed. The plan allows all those involved to carefully evaluate the specific inspection in order to prepare for a successful conclusion. Utilizing Personal Safety concepts prior to a potentially dangerous situation is part of the training programs of many other Federal Agencies. The plan should document what specific roles and responsibilities are needed to conduct the inspection/investigation of sample collection. The plan should also answer the questions: Who, What, Why, When and Where concerning the potential danger.

There are seven principles to a Personal Safety Plan. These are:

1. Summary of Potential Hazards: This section of the personal safety plan includes all of the potential hazards, in a detailed description, that prompted the need for a personal safety plan. Be sure to answer the questions: Who, What, Where, When, and Why. Also include any specific hazards that require personal protective equipment or situations at the facility that may cause allergic reactions for investigators or analysts. Include in the section information from past inspection reports, discussions with previous FDA, State or local investigators, as well as any environmental or plant/facility specific information that would negatively impact a successful personal safety plan when initiated.

2. Sources of Information: This section of the personal safety plan includes all the sources from which your potential hazards were collected. For instance, document which FDA investigator or State inspector supplied factual statements; state the documents or databases from which you obtained information to assist in your hazard summary. This section is important, as it documents factual evidence, similar to all of your other FDA factual inspection gathering information.

3. Response Alternatives: This section will be the most important part of your plan because it includes all of the details of what will be done to mitigate the hazards. In this section, provide a list of factual, practical responses or options to consider. This will also allow your supervisor to see all the possible ways to handle the situation. The response plan should also outline all of the tools that you possess to assist you in handling the situation carefully, including training, experience, and other procedures you have at your disposal. Roles and responsibilities of all involved in the plan should be identified including those intended to be on-site, and those who will be off-site, and participating in the plan.

4. Communication: provide all information about how communication will occur between on-site and off-site participants; between those present on-site, and any emergency, law enforcement or medical responders. Also consider types of communication, e.g. code words for emergencies.

5. Transportation: Provide information in the plan as to how travel to the facility will happen. Is there a coordination point? Do you intend to use Government marked or unmarked cars? Who will ride in each car? What route will be taken going to and leaving the facility? Consider where you will park the car when you arrive at the facility. Consider what modes of
communication will be used to communicate if multiple vehicles are used.

6. Equipment: Include in this section all equipment needed to initiate this plan. Is personal protective equipment needed? Is there any special sampling equipment or other equipment needed? Include in this section, equipment such as communication tools, FDA forms, etc. Assure that the equipment needed is in full functioning mode.

7. Emergency Exit Strategy: Describe in this section what the exit strategy will be in the event of an emergency. Consider emergency strategies for safety (issues), as well as any medical emergency. How will the emergency be communicated on-site and off-site? How do you exit the facility and return to your vehicle? Is there a scheduled meeting point to assure all are safe? The goal is to have no one left behind. Remember to contact your supervisor when you return to safety.

Once the plan has been completed, a debriefing of the situation should occur with all who were involved in the plan development. Evaluate what went well, what needed improvement, what would be done differently the next time. Evaluate whether the plan was successful and document lessons learned for the next time.

The Personal Safety Plan should be developed by the investigator, supervisor, other investigators who may be familiar with the facility, compliance officer, if needed, and any other individuals (District, Region, or HQ experts, etc.) who may be able to assist in the depth, scope, and specifics of the firm in question. The decision of who should be involved in the development and approval of the plan is left to the districts’ discretion.

District management and all involved in writing the personal safety plan should meet when necessary in order to assure a well-developed, and understood personal safety plan. You and your supervisor should maintain contact during the execution of the personal safety plan. The supervisor should contact the employee during these personal safety situations at a predetermined frequency outlined in your plan. A debriefing session should be held following the execution of the plan. Discussions should include what actions worked well and where there are areas of improvement.

For foreign inspections where a Personal Safety Plan is warranted, OO will assist the inspection team. The inspection team’s management may also wish to participate so that there is clear understanding of what actions will be taken for the foreign inspection.

The Personal Safety Plan should be placed in the official establishment file jacket separate from any EIRs in the same location as any Personal Safety Alert memos. A copy of completed and executed Personal Safety Plans must be sent to OO in order for OO to maintain a reference library of all Personal Safety Plans.

5.2.2 - NOTICE OF INSPECTION

Upon arrival at the firm locate the owner, operator or agent in charge of the establishment. This should be the top Management Official on site. Be certain of this individual's status. Introduce yourself by name, title and organization. Show your credentials to this person and present a properly signed, completed original of the FDA 482, Notice of Inspection.

If additional Agency personnel accompany you during the inspection, they must show their credentials to the top Management Official upon arrival at the site. A new FDA 482, Notice of Inspection must be issued. Submit an exact copy of the FDA 482(s) with your EIR. Explain the purpose of your visit. Readily accept any management offer to have a representative accompany you on the inspection.

If non-FDA officials accompany you during your inspection and do not have authority to enter and inspect, you should obtain permission (preferably in advance) from the most responsible individual at the firm. Non-FDA officials and those who do not hold FDA credentials do not sign the FDA 482. See IOM 5.1.1 and 5.10.4.3.3.

For multiple occupancy inspections in drug establishments, refer to IOM 5.1.1.11. Inspections of multiple firms, which are separate legal entities, should be reported under separate EIRs.

If faced with a refusal, or partial refusal of inspection, proceed as outlined in IOM 5.2.5.4.

Any time a FDA 482 is issued, also issue an FDA 484 (at the conclusion of the inspection), Receipt for Samples, if you collect any samples at the firm. See IOM 5.2.4. See IOM 4.1.1.1 and 4.1.1.2 for instructions for issuance of the FDA 482 in certain sampling situations.

If you have concerns of when to or when not to issue the FDA 482, discuss with your supervisor.

5.2.2.1 - Multiple Date Inspections

If your inspection covers more than one day, advise management at the close of each day you have not finished the inspection and when you will return. Do this each day until you finish the inspection. A FDA 482 is not required for each day of an inspection or when different individuals are interviewed. If there will be an extended period of time (i.e., a week or longer) before you can return to the firm to complete the inspection, be sure management is aware of the delay and discuss with your supervisor whether or not you need to issue another FDA 482.

5.2.2.2 - Inspection of Vehicles
If vehicles are present which are owned or leased by the firm being inspected and it is necessary to inspect the vehicles, the inspection of these is covered by the FDA 482, Notice of Inspection, you issued to the firm.

If vehicles (trucks, trailers, RR cars, etc.) which are not owned or leased by the firm are present and inspection is necessary, a separate FDA 482, Notice of Inspection, is required:

1. Issue the FDA 482 to the driver of the vehicle.
2. If the driver is not present and if, after a diligent search, he cannot be located, issue a separate FDA 482 jointly to the firm being inspected and to the firm whose name appears on the cab. Enter the license number of the vehicle on the FDA 482. Give the original FDA 482 to the firm and leave a copy in the cab of the vehicle.
3. If there is no cab present, prepare a separate FDA 482 modified to read "*** to inspect unattended vehicle ***", and issue it to the firm being inspected as the "agent in charge" of the vehicle. Enter the license number of the vehicle, trailer or RR car number, etc., on the FDA 482. Should the firm being inspected refuse to accept the Notice, leave it in a conspicuous place in the vehicle. Describe the circumstances in your EIR.

5.2.2.3 - Follow-Up Inspections by Court Order

At times you may be instructed to conduct inspections of firms by authority of an injunction or other court order. This situation provides separate and distinct inspectional authority involving both the authority of the court order and the authority of Section 704 of the FD&C Act [21 U.S.C. 374], each providing independent courses of action.

When assigned to conduct inspections under these situations, obtain a copy of the injunction or other court order bearing the filing stamp and all relevant signatures. Prior to starting the inspection study the order thoroughly for any special instructions of the court. Your supervisor will assist you in determining the depth of the inspection necessary to cover all of the court requirements.

Take a clearly legible copy of the court decree (not necessarily a certified copy) with you to the firm to be inspected.

Present your credentials in the same manner as for any other EI. Issue the FDA 482, Notice of Inspection, modified to read, "Notice of Inspection is hereby given under authority of injunction (provide here the injunction number and/or other identification) against the firm and pursuant to Section 704 ***". Show the person to whom the FDA 482 was issued a copy of the Order, and, read the following statement to that person.

"This inspection is being conducted under the authority of injunction (add the injunction number and/or other identification) (or other court order) granted by the United States District Court against this firm on (date). The inspection will cover all items specified in the decree. In addition to the inspection authority granted in the court decree, I am issuing you a Notice of Inspection under the authority of Section 704 of the Federal Food, Drug and Cosmetic Act which authorizes inspections of firms subject to that Act."

If the firm refuses access to records, facilities, or information for which the decree provides inspectional authority, read the pertinent section(s) or portion of the order to the person refusing so there will be no misunderstanding as to the requirements of the decree. If the person still refuses, report the facts to your supervisor as soon as possible so the court can be promptly advised of the situation. See IOM 5.2.5 for information on handling refusals.

At the conclusion of the inspection, if a FDA 483 is to be issued and you are using Turbo EIR, follow the Turbo instructions to get injunction specific cites on the FDA 483.

When you prepare your EIR, describe the sequence of events in detail including exactly what happened and how you handled the situation. This documentation will help support any charge of violating the court order and/or Section 704 of the FD&C Act [21 U.S.C. 374].

The court order may require a report to the court. Discuss this with your supervisor since the district will normally handle this part of the requirement.

5.2.2.4 - Conducting Regulatory Inspections When the Agency is Contemplating Taking, or is Taking, Criminal Action

You should not issue a Notice of Inspection if the agency is contemplating taking, or is taking, criminal action against a firm without first discussing the matter with your supervisor. District management will obtain advice from the Office of Chief Counsel and will allow or not allow, the inspection to proceed based on any considerations related to the criminal investigation. Decisions to inspect under such circumstances should be based on considerations of whether or not the request is consistent with FDA's responsibility to assure articles are not produced or distributed in violation of the Federal Food, Drug, and Cosmetic Act or other Federal law within FDA's jurisdiction. The district should ensure these considerations are documented. In no circumstance should an inspection be conducted solely to obtain evidence to support a possible criminal case.

Inspections conducted in accord with this responsibility to protect the public and limited in scope to the authorizing statute are lawful even when criminal action is being considered or pursued. The Fourth Amendment to the United States Constitution prohibits searches without a warrant supported by probable cause. One exception to the warrant requirement includes the inspection of industries long subject to close supervision and inspection, which are conducted under a statute dispenses with the need for a probable cause warrant. Three criteria must be met under this exception from the warrant requirement. First, the regulatory scheme
authorizing the regulatory inspection must be supported by a substantial government interest. Second, regulatory inspections must be necessary to further the regulatory scheme. Third, the statute's inspection program, in terms of the certainty and regularity of its application, must provide a constitutionally adequate substitute for a warrant.

Section 704 of the Federal Food, Drug, and Cosmetic Act [21 U.S.C. 374] is appropriately designed to allow regulatory inspections within appropriate limits. This provides the authority to inspect at reasonable times, within reasonable limits, and in a reasonable manner, establishments or vehicles being used to process, hold, or transport food, drugs, devices, or cosmetics. See IOM 2.2.1.1. FDA's normal inspection procedures provide guidance on what should be considered reasonable under Section 704.

If you become aware of an ongoing criminal investigation, notify your supervisor. The district should follow the Regulatory Procedures Manual (RPM) and notify the appropriate Center of any OCI involvement in a Center directed inspection.

The discovery of evidence of a criminal violation may also be relevant to FDA's responsibility to assure articles are being produced in conformity with the Food, Drug, and Cosmetic Act. Additional inspections may be warranted. Such inspections should be planned and documented in accordance with the preceding section, "Conducting Regulatory Inspections When the Agency is Contemplating Taking, or is Taking, Criminal Action."

5.2.2.6 - Use of Evidence Gathered in the Course of a Criminal Investigation

The extent to which information gathered in the course of a criminal investigation may be shared with other components of FDA will vary with each case. Investigators should determine the extent of information sharing in accordance with the following guidelines.

Information and evidence gathered in the course of a criminal investigation may be shared with regulatory personnel, subject to two reservations:

1. Information obtained pursuant to grand jury subpoena or testimony may not be shared. Disclosure of such information to anyone other than individuals identified by the Department of Justice attorney involved could subject the individual making the improper disclosure to sanctions for contempt by the court. Only the court can authorize disclosure beyond these parameters. Information obtained by other means (search warrant, cooperative witnesses, surveillance, etc.) may be shared, subject to the following paragraph.

2. There may be a need to protect the confidentiality of the criminal investigation. For example, disclosure to regulatory investigators might prematurely disclose the existence of the criminal investigation or the identity of confidential informants. However, whenever you are calculating the need to protect the confidentiality of information gathered in the course of a criminal investigation through means other than the grand jury, you must consider whether it will be in the interest of public health to protect the confidentiality of that information.

Criminal investigators should consult their supervisors to determine whether disclosure should be made to regulatory investigators.

5.2.2.7 - Use of Evidence Voluntarily Provided to the Agency

Criminal and regulatory investigators may share information and evidence voluntarily provided to FDA, without use of the regulatory inspection authority, search
warrant, or subpoena. If criminal investigators decide not to share such information because of a need to protect the confidentiality of the criminal investigation, they should consider the potential impact on the public health of protecting the confidentiality of that information.

5.2.2.8 - Concurrent Administrative, Civil, and Criminal Actions

It may be appropriate to seek administrative and/or civil remedies against a firm or individual under investigation for criminal violations. There are many issues involved in determining whether such actions may proceed concurrently, or whether certain actions should proceed first. Each situation must be evaluated on an individual basis. If administrative and/or civil remedies are under consideration against a firm or individual also under investigation for criminal violations, representatives from the Center responsible for evaluating the administrative and/or regulatory action should meet with the Office of Criminal Investigations Headquarters staff to discuss issues related to the timing of administrative, civil, and criminal actions. The Office of Criminal Investigations and other components of FDA may share information subject to the reservations set out earlier.

5.2.2.9 - Working with a Grand Jury

Finally, if you are assigned to work with a grand jury, you should not participate in a regulatory inspection or other regulatory matter involving the same firm or individual(s). Such participation is contrary to long standing agency policy, might be unlawful, and could result in sanctions against the investigator and the agency. You should not participate in any regulatory matters that could result in improper disclosure of grand jury information, even after the grand jury investigation is closed. Grand jury proceedings remain secret even after they are concluded. Under no circumstances should you undertake such participation without first obtaining clearance from the Department of Justice attorney or the Office of Chief Counsel attorney assigned to the grand jury case. See IOM 2.2.7.3 for additional information on Grand Jury proceedings.

5.2.3 - REPORTS OF OBSERVATIONS

The FDA 483, Inspectional Observations (see Exhibit 5-5) is intended for use in notifying the inspected establishment’s top management in writing of significant objectionable conditions, relating to products and/or processes, or other violations of the FD&C Act and related Acts (see IOM 5.2.3.2) which were observed during the inspection. These observations are made when in the investigator’s “judgment”, conditions or practices observed, indicate that any food, drug, device, or cosmetic have been adulterated or are being prepared, packed, or held under conditions whereby they may become adulterated or rendered injurious to health. The issuance of written inspecational observations is mandated by law and ORA policy.

All FDA-483s should adhere to the following general principles:

1. Observations which are listed should be significant and correlate to regulated products or processes being inspected.
2. Observations of questionable significance should not be listed on the FDA-483, but will be discussed with the firm’s management so that they understand how uncorrected problems could become a violation. This discussion will be detailed in the EIR.

All FDA-483s should have the following characteristics to be useful and credible documents:

1. Each observation should be clear and specific.
2. Each should be significant. Length is not necessarily synonymous with significance.
3. Observations should not be repetitious.
4. The observations should be ranked in order of significance.
5. All copies of the FDA-483 should be legible.

If an observation made during a prior inspection has not been corrected or is a recurring observation, it is appropriate to note this on the FDA 483.

As of 1997, ORA established a FDA 483 annotation policy for medical device inspections. See IOM 5.2.3.4.

Regardless of whether an establishment’s FDA 483 is annotated, investigators and analysts should make every reasonable effort to discuss all observations with the management of the establishment as they are observed, or on a daily basis, to minimize surprises, errors, and misunderstandings when the FDA 483 is issued. This discussion should include those observations, which may be written on the FDA 483 and those that will only be discussed with management during the closeout meeting. Industry may use this opportunity to ask questions about the observations, request clarification, and inform the inspection team what corrections have been or will be made during the inspection process. Investigators are encouraged to verify the establishment’s completed corrective actions as long as the verification does not unreasonably extend the duration of the inspection.

As of April 2010, ORA established a policy of reporting positive environmental samples on the FDA 483 prior to closeout for food inspections. This policy is applicable only if the results are known before conclusion of the inspection; the investigator should not prolong the inspection to include the results.

There may be instances where same day discussion of observations may not be possible due to the volume of documents collected and document review reveals observations on a different day than the documents were collected or in other circumstances. When these instances occur immediately prior to the conclusion of the inspection
the lack of a daily discussion of observations does not preclude listing of significant observations which were not previously discussed on the FDA 483.

Turbo EIR

Turbo EIR is an automated FDA 483 and EIR reporting system. Use Turbo EIR to generate the FDA 483 where applicable cite modules exist. Turbo EIR should not be used to create a FDA 483 during an inspection of a firm involving multiple commodity areas when FDA 483 cites do not exist for ALL of the commodity areas for which observations need to be included on the FDA 483. You should be able to write the entire FDA 483 using Turbo EIR.

Use Turbo EIR for all EIRs whether or not your FDA 483 was generated using Turbo and when no FDA 483 was issued. See IOM 5.10.4.

5.2.3.1 - Preparation of Form FDA 483

It is not necessary to complete all headings of the FDA 483, when multiple page 483s are issued. Complete all headings on the first page and, on subsequent pages, only those necessary to identify the firm and dates inspected. FDA 483s should be issued at the conclusion of the inspection and prior to leaving the premises. However, in preparing some complex FDA 483s, it may be necessary to leave the premises and return at a later time to issue and discuss your inspectional observations. In this case, you should advise the firm’s management your inspection has not been completed and you will return to issue the FDA 483 and discuss inspectional findings. There should be no unreasonable or unwarranted delays in issuing and discussing the FDA 483. During the inspection, do not show the firm’s management a draft, unsigned copy of the FDA 483 or an electronic copy of the FDA 483 on your computer screen. You should issue only those necessary to identify the firm and dates inspected. FDA 483s should be issued at the conclusion of an inspection need not prevent issuance of the FDA 483. However, absence of a team member at the conclusion of an inspection need not prevent issuance of the FDA 483. See IOM 5.1.2.5.1. If you use an electronically generated FDA 483, assure you have a copy for the District files -- an unsigned photocopy or printed duplicate is unacceptable. See IOM 5.2.3.6.2.

5.2.3.1.2 - SIGNATURE POLICY

Everyone present at issuance signs the first and last pages of the FDA 483 and initials each intervening page in the signature block.

Note: if you are not using the official multi-part FDA 483 form and a copier is not available, insert carbon paper to reproduce a signed copy of the FDA 483.

See IOM 5.2.3.6-Distribution of the FDA 483.

5.2.3.1.3 - DATE ISSUED

Enter the date the form is actually issued to the firm’s management.

5.2.3.1.4 - OBSERVATIONS

“During an inspection of your firm (I) (We) observed” - Where applicable, when formulating each FDA 483 observation, answer Who (using titles or initials when necessary), What, When, Where, How, and challenge each observation by asking So What? (regarding its significance)

Enter your reportable observations succinctly and clearly. Conditions listed should be significant and relate to an observed or potential problem with the facility, equipment, processes, controls, products, employee practices, or records. “Potential problems” should have a reasonable likelihood of occurring based upon observed conditions or events. Do not cite deviations from policy or guidance documents on your FDA 483.

As appropriate, FDA 483 observations should include relationship of observations to a given population, for example, “Two out of 50 records examined were * * *” or “4 out of 12 bags examined were ***.” When appropriate, a FDA 483 observation may refer to inadequate situations
as long as you provide supporting facts (examples) or explanation as to why the condition, practice or procedure observed is inadequate.

It is preferred not to identify individuals or firms by name i.e., suppliers and consignees within the FDA 483. Where appropriate to support the FDA 483 observation, identify the individual(s) or firm(s) by substituting other non-specific identifying information as below. Document your evidence in your EIR, fully explaining the relationship(s).

1. The lot number for a component received from or shipped to firm "A".
2. The invoice number for a shipment from or to firm "A".
3. A patient #, record #. See IOM 5.2.3.3 item 7.
4. The study number for a particular Clinical Investigator site.
5. Other necessary but non-specific identifying information to show the observation's relationship to a particular firm and/or individual.

Presently there are three ways to issue a FDA 483.
1. Turbo EIR Field Agent
2. Traditional hard copy FDA 483.
3. Electronic (non-turbo EIR) version of the FDA 483.

When using a traditional hard copy FDA 483 or electronic (non-turbo) version of the FDA 483, the current version of the 483 must be used. As of the printing of the 2014 IOM, the current version of the FDA 483 is dated 9/08.

5.2.3.1.5 - MEDICAL DEVICE INSPECTIONS

The following language should be inserted on the FDA 483 in addition to the above statement: "The observations noted in this form FDA 483 are not an exhaustive listing of objectionable conditions. Under the law, your firm is responsible for conducting internal self-audits to identify and correct any and all violations of the quality system requirements."

5.2.3.1.6 - CORRECTION OF FDA 483 ERRORS

These procedures do not pertain to adverse conditions noted and then corrected during the inspection. Observations of this type stand and should remain on the FDA 483.

The Inspectional Observations (FDA 483) is of critical importance to both the Agency and regulated industry. Individual FDA 483s may become public through publishing in industry trade press, FOI inquiries, Headquarters postings and other means. Therefore, complete and accurate documentation of corrections to this official document is critical.

5.2.3.1.6.1 - Errors Discovered Prior to Leaving the Establishment

Non-Turbo, FDA 483s:

1. Make handwritten changes to correct the error/s on the original FDA 483 and initial the changes. Correct errors by striking through the erroneous text and entering the correct information (if any). When possible retrieve and destroy all uncorrected copies of the FDA 483 either provided to or produced by the establishment.
2. If the establishment has photocopying equipment available and will provide you with a copy of the corrected original FDA 483 then obtain a copy of the corrected original document from the establishment. If the establishment has no such equipment or refuses to provide you with a copy of the original corrected FDA 483 then make the corrections and initial the changes using carbon paper and retain the carbon copy of the corrected FDA 483 for your District's official establishment file.

Turbo FDA 483s - All corrections/deletions should be made in Turbo. If there are technical difficulties which prevent you from issuing a modified Turbo 483, you may handwrite the corrections on the original (maintain a copy for the EIR) and inform the firm representatives that you will make corrections and provide them with the corrected Turbo 483.

1. Changes made to correct errors in the text of the observation will show on the face of the final printed FDA 483. Changed Text deletions will remain visible as strike through and correction made. For example, "lot 4234 5678" – (select text, right click, select font and select strike-through) or from "lot 1234" to "lots 1234 and 5678" and bold the changes "lots 1234 and 5678".
2. If an entire observation is removed or the underlying citation is changed, incidental text will be used to add the statement "An observation concerning *** was removed [or the underlying citation was changed] based on discussions with management."
3. Addition of a new observation or changes to the observation.

5.2.3.1.6.2 - Errors Discovered after Leaving the Establishment

Normally, you should not use addenda/amendments to issue additional FDA 483 items after the inspection has been closed out and you have left the premises.

1. Non-Turbo, FDA 483s: Discuss any errors with your supervisor. If necessary a revised FDA 483 will be prepared.
2. Turbo FDA 483s: Discuss any errors with your supervisor. Make all corrections/deletions in Turbo. Changes made to correct errors in the text of the observation will show on the face of the final printed FDA 483. Changed Text deletions will remain visible as strike through and additions added.
3. Issuing FDA 483s: Personally deliver the revised and/or corrected FDA 483 addendum/amendment to the firm for discussion. If personal delivery is not practical, mail the addendum/amendment to the firm with a full explanation cover letter. Include a copy of the original FDA 483, and a copy of the letter in the EIR. In addition, you should call the person to whom
the original FDA 483 was issued, to discuss the change(s). Document your discussion in your EIR.

NOTE: The issuance of an amended FDA 483 in person or via mail does not change the inspectional end date. The inspectional end date remains as the date the original FDA 483 was issued.

5.2.3.2 - Reportable Observations

You should cite factual observations of significant deviations from the FD&C Act [21 U.S.C. 301], PHS Act, 21 CFR, and other acts where FDA has enforcement authority unless these cites require concurrence or are specifically prohibited – see IOM 5.2.3.3 Non-Reportable Observations. Examples of these observations generally fall into two categories.

5.2.3.2.1 – ADULTERATION OBSERVATIONS

Review Sections 402, 501, 505(k), 601, and 704 of the FD&C Act [21 U.S.C. 342, 351, 355(k), 361, and 374]. Include specific factual observations of:
1. Foods, drugs, devices, or cosmetics consisting in whole or in part of filthy, putrid, or decomposed substances.
2. Undesirable conditions or practices, bearing on filth or decomposition, which may reasonably result in the food, drug, device, or cosmetic becoming contaminated with filth.
3. Insanitary conditions or practices which may reasonably render the food, drug, device, or cosmetic injurious to health.
4. Careless handling of rodenticides or pesticides.
5. Results of field tests (organoleptic examination of fish, crackout of nuts, etc.) if the results revealed adulteration.
6. Observations of faulty manufacturing, processing, packaging, or holding, of food, drug, or device products as related to current good manufacturing practice regulations including inadequate or faulty record keeping.
7. Observations of faulty can closures and/or deviations from recommended processing times and temperatures.
9. Results of analytical laboratory findings which reveal adulteration.

5.2.3.2.2 - OTHER OBSERVATIONS

You may include other factual observations of significant deviations from the FD&C Act [21 U.S.C. 301], 21 CFR, Government Wide Quality Assurance Program (GWQAP) requirements, and other Acts as directed by CPs and other agency directives. In some cases, you may cite labeling deviations as directed below. This list is not all inclusive.

2. Observations, forming the basis for product non-acceptance under the Government Wide Quality Assurance Program (GWQAP). See IOM 5.2.3.5.
3. Deviations from blood and blood products labeling requirements as specified in 21 CFR 606.121 and 21 CFR 640.
4. Animal protein products, and feeds containing such products, that are not in compliance with the labeling requirements of paragraphs (c) through (f) of 21 CFR 589.2000. See Section 403(a)(1) or 403(f) of the FD&C Act [21 U.S.C. 343(a)(1) or 343(f)].
5. Deviations from the applicable labeling regulations for human cells, tissue, and cellular and tissue-based products (HCT/Ps) as specified in 21 CFR 1271 and CP 7341.002.
6. Observations indicating drug misuse, failure to maintain proper drug use records, and/or poor animal husbandry practices during tissue residue investigations. See the applicable Compliance Program(s) for guidance.
7. Observations indicating non-conformity with the postmarketing adverse drug experience reporting requirements as specified in 21 CFR 310.305, 314.80, 314.98, 315.30, and 600.80 or other postmarketing requirements as specified in 21 CFR 314.81 or 600.14. See Sections 506 and 760 of the FD&C Act [21 U.S.C. 355(k) and 379aa].
8. Observations indicating non-conformity with the Medical Device Reporting requirements as specified in 21 CFR 803 [See Section 519(a) of the FD&C Act [21 U.S.C. 360(i)]; the Medical Devices Reports of Corruptions and Removals requirements as specified in 21 CFR 806 [See Section 519(f) of the FD&C Act [21 U.S.C. 360(if)]; and the Medical Device Tracking requirements as specified in 21 CFR 821 (See Section 519(e) of the FD&C Act [21 U.S.C. 360(e)]).
9. Observations indicating non-compliance with medical device pre-market notification requirements and pre-market approval requirement under FD&C Act sections 510(k) and 515 [21 U.S.C. 360(k) and 360e] respectively, should only be made with the prior confirmation of CDRH and/or CBER. 21 CFR PART 200.10 does allow reporting observations noted at a contract facility to the contracting facility. Before doing this, check with your supervisor to determine if this is appropriate.
10. Observations indicating non-compliance with LACF/Acidified food registration and failure to file scheduled processes. Before doing this, verify lack of such, as covered in CP 7303.803A.

5.2.3.3 - Non-Reportable Observations

Do not report opinions, conclusions, or characterize conditions as "violative." The determination of whether any condition is violative is an agency decision made after
considering all circumstances, facts and evidence. See IOM 5.2.7 involving discussions with management at which time opinions may be discussed.

Do not quote Regulations (e.g., specific CFR sections) when listing items.

Do not report observations pertaining to:
1. Label and labeling content, except per IOM 5.2.3.2.2, items 2, 3, 4 and 5 above.
2. Promotional materials.
3. The classification of a cosmetic or device as a drug.
4. The classification of a drug as a new drug.
5. Non-conformance with the New Drug Regulations, 21 CFR 312.1 (New Drugs for Investigational Use in Human Beings: Exemptions from Section 505(a)) unless instructed by the particular program or assignment.
6. The lack of registration required by Section 415 and 510 of the FD&C Act. The lack of registration per 21 CFR 1271 Subpart B Procedures for Registration and Listing, promulgated under Section 361 of the PHS Act.
7. Patient names, donor names, etc. If such identification is necessary, use initials, code numbers, record numbers, etc.
8. Corrective actions. Specific actions taken by the firm in response to observations noted on the FDA 483 or during the inspection are not listed on the FDA 483, but are reported in the EIR. Except as described in IOM 5.2.3.4.
9. The use of an unsafe food additive or color additive in a food product.

Use Turbo EIR to document in the “General Discussion with Management” section Non-Reportable Observations, which you discussed with management. These objectionable conditions fall into three basic categories:
1. Observations of significant deviations from specific Laws and/or regulations, non-reportable items 1-9 above.
2. Observations of deviations from specific Laws and/or regulations, which in your judgment, are of “questionable significance” and “deemed not to merit inclusion on the FDA 483,” but do warrant discussion with management.
3. Observations which in your judgment deviate from official published guidance, not regulations, but warrant discussion with management.

The reporting of observations in these 3 categories is as follows:

Category 1: You should select the appropriate Turbo cite, verify or set the “Print type” to “Do Not Print,” and save the observation in the Turbo database. This should be done even if there are no other reportable observations. For example, Lack of Food Registration as covered in IOM 5.4.1.5.3 is not reportable.

Category 2 or 3: You should always report these two categories of observations which were discussed with management under the “General Discussion with Management” heading in the EIR as specified by IOM 5.10.4.3.15. You have options in choosing how observations in category 2 are reported. You may select the appropriate cite in Turbo, enter the “specifically” text regarding the observation, and discussion with management, set it to “Do not print”, save, and it will be automatically entered into the Turbo EIR when it is generated.

The second option which is also true for category 3 (i.e., there are no Turbo cites for official guidance, only regulations) is the observation/s discussed with management may be entered directly into the Turbo EIR under the “General Discussion with Management.”

5.2.3.4 - Annotation of the FDA 483

Offer to annotate the FDA 483 for all medical device inspections. The district has discretion to annotate the FDA 483s in other program areas. BIMO inspections are generally excluded from annotations. Annotations of FDA 483s for inspections in other program areas may be done if both the establishment and the investigator/team believe annotation will facilitate the inspection process. When a FDA 483 is annotated it should be done in accordance with the guidance that follows.

Inform the establishment of the annotation program at some point prior to the final discussion with management. Determine from management whether they wish to have their FDA 483 observations annotated. It is voluntary on the part of the establishment. If the establishment does not want one or more observations annotated, you must honor the request.

The actual annotation of the FDA 483 should occur during the final discussion with management. The annotations are succinct comments about the status of the FDA 483 item. It is not permissible to pre-print or pre-format the annotations onto the FDA 483 form. The annotations can be made after each observation, at the end of each page of the FDA 483 or at the bottom of the last page of the FDA 483 prior to the investigator's signature. The establishment should review the annotations on the issued FDA 483 to ensure there are no misunderstandings about promised corrective actions. See IOM 5.2.3 for discussions of FDA 483 observations with management.

If the establishment has promised and/or completed a corrective action to an FDA 483 observation prior to the completion of the inspection, the FDA 483 should be annotated with one or more of the following comments, as appropriate:
1. Reported corrected, not verified.
2. Corrected and verified.
3. Promised to correct.
4. Under consideration.

The term "verified" means "to confirm; to establish the truth or accuracy". In this case, you must do the verification. In some situations, you will not be able to
verify the corrective action unless there is further district or Center review or until there is another inspection of the establishment.

The establishment's stated objections to any given observation or to the FDA 483, as a whole should not be annotated on the FDA 483. If they would prefer no annotation, do not annotate it. The EIR should include the establishment's objections to the observation and the fact the establishment declined to have the observation annotated.

When an establishment has promised corrections and furnishes a date or timeframe (without a specific date) for completion, then you may add "by xxx date" or "within xxxx days or months" in the annotation. Where the investigator and the establishment have "agreed to disagree" about the validity of an observation, you may annotate this observation with "Under consideration" or with no annotation based on the establishment's desire.

All corrective actions taken by the establishment and verified by FDA should be discussed in detail in the Establishment Inspection Report (EIR) and reported using the Compliance Achievement Reporting Systems (CARS).

5.2.3.5 - Government Wide Quality Assurance Program (GWQAP)

When performing product acceptance examinations under the GWQAP, you must discuss all deficiencies with management and report these deficiencies in writing on the FDA 483. This includes all deficiencies related to the FD&C Act as well as deficiencies in complying with contract requirements, which result in non-acceptance. There must be a clear differentiation on the FDA 483 between these two types of deficiencies.

Enter the FD&C type deficiencies (GMP deviations, etc.) first on the FDA 483. If there are deficiencies in contract provisions, draw a line across the sheet and add a heading "The Following Additional Contract Non-Conformances Were Observed." Enter each deficiency, which forms a basis for non-acceptance, followed by the reference to the applicable contract requirement or specification.

5.2.3.6 - Distribution of the FDA 483

Be sure all copies of the original FDA 483 are legible and distribute as follows.

5.2.3.6.1 - ORIGINAL

The FDA 483 issued to the firm signed in pen and ink.

Before leaving the premises at the end of the EI present the original to the individual who received the FDA 482, Notice of Inspection, if the person is present and qualifies as "most responsible." If the person is not available or is outranked by someone else, present it to the individual who meets the definition of owner, operator, or agent in charge.

5.2.3.6.2 - COPIES

Replicas of the "original".

Attach one copy of all FDA 483s issued to the firm to the EIR. This includes turbo or non-turbo copies of any signed, modified, and/or amended FDA 483, or 483 addenda. See IOM 5.2.3.1.6 (Correction of FDA 483 Errors). A copy may be sent to the top management of the firm including foreign management, unless the individual to whom you gave the original is the top official of the firm.

If the inspection covered vehicles as described in IOM 5.2.2.2, leave an exact copy of the list of observations with the firm being inspected. The original will be sent by your district to the firm owning or leasing the vehicle. You must make every effort to obtain the name and address of the vehicle owner. Usually the firm name is on the vehicle; however, it may require a trace of the vehicle license number. Discuss with your supervisor before taking this step. See IOM 4.4.7.2.

5.2.4 - RECEIPT - FACTORY SAMPLES

You must issue an FDA 484, Receipt for Samples, if you collect any physical sample during an inspection. At the end of the EI and prior to leaving the premises, issue the original FDA 484 to the same individual who received the FDA 482. (See IOM 4.2.5) If this person is not available, give it to someone else who meets the definition of owner, operator, or agent in charge. Submit an exact copy with the EIR. Do not comment on type of examination expected or promise a report of analysis.

5.2.4.1 - Items Requiring Receipt

Issue a FDA 484 for any item of food, drug, device, or cosmetic actually removed from the establishment. NOTE: A receipt must always be issued to anyone from whom you obtain Rx drugs. This includes individuals as well as firms. See IOM 4.2.5.4 and IOM 4.4.10.3.44.

The following are examples of exhibit materials also requiring a Receipt for Samples:
1. Air filter pads,
2. Rodent pellets, and
3. Any other physical evidence actually removed from the plant.

5.2.4.2 - Items Not Requiring Receipt

Do not issue a FDA 484 for:
1. Items or materials examined during the inspection but not removed from the establishment (report adverse results of analysis of materials on FDA 483 as indicated in IOM 5.2.3.2),
2. Labels or promotional material,
3. Photographs taken during the inspection,
4. Record(s): including production, quality control, shipping and interstate records.

Firm management may request copies of documents or records you obtain from their firm. There is no objection to supplying them.

See IOM 5.3.8.5 for procedures when a firm requests a receipt for records copied during an inspection or investigation.

5.2.5 - INSPECTION REFUSAL

Refusal as used in your IOM means, refusing to permit an inspection or prohibiting you from obtaining information to which FDA is entitled under the law. See IOM 4.2.3 for information regarding refusal to permit sampling.

In the case of a refusal you must show your conduct was reasonable, fair, and you exercised reasonable precaution to avoid refusal. You must have shown your credentials and given the responsible individual a properly prepared and signed Notice of Inspection, FDA 482.

Inspection refusals may take several forms. All refusals to permit inspection must be reported in your EIR under the "Refusals" heading.

In the case of drug inspections, inspection refusals, as well as delaying, denying, or limiting your ability to conduct the inspection, may cause a drug to be deemed adulterated under Section 501(j) of the FD&C Act [21 U.S.C. 351(j)]. See subsection 5.5.5.8 in Subchapter 5.5 (Drugs) for further guidance on responding to these situations.

5.2.5.1 - Refusal of Entry

When you are faced with a refusal of entry, call the person’s attention to the pertinent sections of the Acts (Sections 301(f) and 704 of the FD&C Act [21 U.S.C. 331 (f) and 374] and Section 351(c), 360A(a), (b) and (f); 360B(a); and 361(a) of the Public Health Service Act. Portions of these are listed on the front and back of the FDA 482. If entry is still refused, leave the completed FDA 482, leave the premises and telephone your supervisor immediately for instructions.

In the case of drug inspections, if the person refuses entry or delays, denies, or limits your ability to conduct the inspection, also call the person’s attention to Section 501(j) of the FD&C Act [21 U.S.C. 351(j)] (an adulterated drug could lead to further prohibited acts under 301(a), (b), (c) [21 U.S.C. 331(a), (b), (c)]).

Furthermore, if during a drug inspection management refuses access to or copying of any record to which you are entitled under law, without giving a reasonable explanation such as requiring sufficient time to compile a large volume of records or translate the records into English, you may call their attention to 501(j) of the FD&C Act. Similarly, if management limits your access to or ability to copy any record to which you are entitled under law, you may call their attention to Section 501(j) of the FD&C Act. See subsection 5.5.5.8 in Subchapter 5.5 (Drugs) for further guidance on responding to these situations.

5.2.5.2 - Refusal to Permit Access to or Copying of Records

If management objects to the manner of the inspection or coverage of specific areas or processes, do not argue the matter but proceed with the inspection. However, if management refuses to permit access to or copying of any record to which you are entitled under law, call attention to Section 301(e) of the FD&C Act [21 U.S.C. 331] or applicable sections of the PHS Act. If management still refuses, proceed with the inspection until finished.

In the case of drug inspections, if management refuses access to or copying of any record to which you are entitled under law, in addition to Section 301(e) noted above, call attention to Section 501(j) of the FD&C Act [21 U.S.C. 351(j)] (an adulterated drug could lead to prohibited acts under 301(a), (b), (c) [21 U.S.C. 331(a), (b), (c)]).

5.2.5.3 - Refusal after Serving Warrant

If you have been refused entry, obtained a warrant, tried to serve or execute it and are refused entry under the warrant, inform the person, the warrant is a court order and such refusal may constitute contempt of court. If the warrant is not then immediately honored (entry and inspection permitted), leave the premises and promptly telephone the facts to your supervisor.

If you have served the warrant and during the inspection you encounter partial refusal or resistance in obtaining access to anything FDA is authorized to inspect by the warrant, inform the firm that aspect of the inspection is part of a court order and refusal may constitute contempt of court. If the warrant is not then immediately honored, leave the premises and promptly telephone the facts to your supervisor.

5.2.5.4 - Hostile and Uncooperative Interviewees

More often than not, investigations or inspections are conducted in a reasonable atmosphere. Nonetheless, there will be times you are confronted by unfriendly or hostile persons.
Your activities must always be conducted with tact, honesty, diplomacy, and persuasiveness. Even though you must at times adopt a firm posture, do not resort to threats, intimidation, or strong-arm tactics.

Many times a hostile or uncooperative attitude on the part of individuals being interviewed results from fear, timidity, or previously distasteful encounters with law enforcement personnel. In most cases a calm, patient, understanding and persuasive attitude on your part will overcome the person's reluctance or hostility. Often the mere fact you patiently listen while individuals share their views will make them receptive to your quest.

5.2.5.4.1 - INDICATORS

Normally you have no way to predict the nature of the individuals you meet. However, there are often indicators, which can alert you, such as:

1. Establishment inspection reports, endorsements or memorandums may show situations where investigators encountered belligerent or hostile individuals. These reports may be FDA reports and/or State contract reports, if available.
2. Discussions and conversations with FDA, federal, state and local inspectors and investigators may reveal instances where uncooperative individuals and problem situations were encountered.
3. The nature of the assignment, program or information requested may indicate some degree of caution is needed.
4. A firm located in an area with a reputation for unfriendliness to law enforcement personnel should alert you some employees of the firm may be less than cooperative during the investigation.

If you find yourself in a situation which, in your judgment, indicates violence is imminent, stop the operation and make an exit as soon as possible. Immediately report the facts to your supervisor.

5.2.5.4.2 - SAFETY PRECAUTIONS

The FDA recognizes there are situations where it is advisable to take precautions for your personal safety. In those, consult your supervisor. Some procedures, which may be utilized to minimize the danger, include:

1. Inspections or investigations carried out by a team of two or more persons.
2. Consider whether or not the use of an unmarked government car would be more beneficial to assist you in your inspection in lieu of a marked government car.
3. Request additional information from your State and/or Local Agencies who also regulate and inspect the facilities in question. In many instances, your State counterparts may have more information regarding the facility. This may be especially helpful for those firms that FDA has not yet inspected but were inspected by your State counterparts.
4. Each government car or inspection team should be assigned one FDA cell phone or alternate communication device. While we recognize that some investigators carry a personal cell phone, FDA strongly suggests that your personal cell phone not be utilized to contact the firm or firm’s management. In some instances, such uses in the past have resulted in later inappropriate contacts from the firm to the individual FDA investigator.

5. Request assistance from local law enforcement agencies prior to or during investigations. This assistance may include information about the facility you are to inspect, assistance with communication devices, or police protection, if the police jurisdiction allows for such an action.
6. In potentially hazardous investigations such as methadone or schedule II Class Drugs, two investigators may be used and personnel from the U.S. Drug Enforcement Administration, State, or local law enforcement agencies may be requested to accompany you.

5.2.5.4.3 - PROCEDURES WHEN THREATENED OR ASSAULTED

In instances when you are actually assaulted or threatened, you should immediately notify your supervisor. Your supervisor can summon local police, United States Marshals, or contact OCI headquarters for assistance (301-294-4030). OCI can make contacts with local police and federal agencies based on previous liaison. Also, the District should notify the Office of Operations via e-mail ORAHQOMPTOCSOSAF@FDA.HHS.GOV.

If you are physically attacked, you have the same recourse as any other citizen as well as the benefit of federal laws protecting government officials while in the performance of their official duties. If you are physically attacked, you should get to safety, call your supervisor, report the incident and seek medical attention if needed. Remember that the medical attention you receive may be used as documentation for the Agency in support of any legal action taken against the firm or the individual.

5.2.5.4.4 - NOTIFICATION OF FBI AND US ATTORNEY

It is a federal crime for anyone to kill, assault, resist, oppose, impede, intimidate, or interfere with, a federal official in the performance of their official duties.

In case of assault or threat against you, notify your supervisor immediately, so the facts can be submitted to the Federal Bureau of Investigations and the U.S. Attorney's office for immediate action.

The referenced sections in Title 18 of the U.S. Code are:

1. Title 18 U.S.C.A. Section 111, which provides: "111. Assaulting, resisting, or impeding certain officers or employees. Whoever forcibly assaults, resists, opposes, impedes, intimidates, or interferes with any person designated in Section 1114 of this title while engaged in or on account of the performance of his official duties, shall be fined not more than $5,000 or imprisoned not more than three years, or both.

Whoever, in the commission of any such acts uses a deadly or dangerous weapon, shall be fined not more

Title 18 U.S.C.A. Section 1114, which provides: "1114. Assisting, protecting, or calling for the aid of another officer or employee. Whoever, in the commission of any such acts uses a deadly or dangerous weapon, shall be fined not more than $5,000 or imprisoned not more than three years, or both.

Whoever, in the commission of any such acts uses a deadly or dangerous weapon, shall be fined not more than $5,000 or imprisoned not more than three years, or both.

Whoever, in the commission of any such acts uses a deadly or dangerous weapon, shall be fined not more than $5,000 or imprisoned not more than three years, or both.

Whoever, in the commission of any such acts uses a deadly or dangerous weapon, shall be fined not more than $5,000 or imprisoned not more than three years, or both.

Whoever, in the commission of any such acts uses a deadly or dangerous weapon, shall be fined not more than $5,000 or imprisoned not more than three years, or both.

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Whoever, in the commission of any such acts uses a deadly or dangerous weapon, shall be fined not more than $5,000 or imprisoned not more than three years, or both.

Whoever, in the commission of any such acts uses a deadly or dangerous weapon, shall be fined not more than $5,000 or imprisoned not more than three years, or both.
that individual the original signed Inspection Warrant, should be made promptly and usually no later than 10 days following its execution.

5.2.7 - DISCUSSIONS WITH MANAGEMENT

After completion of the inspection, meet with the highest ranking management official possible to discuss your findings and observations. The FDA 483 is not a substitute for such discussion since there may be additional questionable practices or areas not appropriate for listing on this form.

During the discussion be frank, courteous and responsive with management. Point out the observations listed on the FDA 483, are your observations of objectionable conditions found during the inspection, and explain the significance of each. Try to relate each listed condition to the applicable sections of the laws and regulations administered by the FDA.

If significant deviations are observed during the inspection, you should inform management during the closeout discussion, the conditions observed may, after further review by the Agency, be considered to be violations of the Food, Drug and Cosmetic Act or other statutes. Legal sanctions available to FDA may include seizure, injunction, civil money penalties and prosecution.

Do not be overbearing or arbitrary in your attitude or actions. Do not argue if management voices a different view of the FDA 483 observations, or of your opinions. Explain, in your judgment the conditions you observed MAY be determined by the FDA, after review of all the facts, to be violations. Make clear the prime purpose of the discussion is to call attention to objectionable practices or conditions, which should be corrected.

Obtain management's intentions regarding correcting objectionable conditions. They may propose corrections or procedural changes and ask you if this is satisfactory. If this involves areas where your knowledge, skill, and experience are such that you know it will be satisfactory, you can so advise management. Do not assume the role of an authoritative consultant. In areas where there is any doubt, you must explain to management you cannot endorse the proposed corrections. Advise the individuals their firm's response may impact FDA's determination of the need for follow-up action, if FDA receives an adequate response to the FDA 483 within 15 business days of the end date of the inspection. A record of the firm's response should be entered into FACTS 483 Firm Response screen.

FDA will supply comments (see RPM 4-1-3 #4) if the establishment will submit its request and its proposed corrections or procedures in writing to the district office.

Concentrate on what needs to be done rather than how to do it. Do not recommend the product or services of a particular establishment. If asked to suggest a product or
consulting laboratory, refer the inquirer to a classified directory or trade publications and or organizations.

Report in your EIR all significant conversations with management or management representatives. In most instances it is not necessary to quote management’s response verbatim. Paraphrasing the replies is sufficient. However, if the situation is such that quoting the reply or replies is necessary, enclose them in quotation marks.

### 5.2.7.1 - Protection of Privileged Information

You have certain responsibilities under the FD&C Act, Section 301(j); Sections 359(d) and 306(e) of the Public Health Service Act; and Section 1905 of the Federal Confidential Statute (18 U.S.C. 1905) regarding protection of confidential material obtained during your official duties. See IOM 1.4.

Do not volunteer information about other firms or their practices. Ignore casual exploratory questions or remarks from management about competitors or their processes. Your casual and seemingly innocuous remarks may reveal privileged information. Therefore, be alert and avoid voluntarily or unknowingly divulging information, which may be privileged or confidential and possibly compromise FDA’s and your own integrity.

Management often request copies of any documents or records you obtain from their firm. There is no objection to your supplying these. When management requests copies of photos taken by you in a plant, follow IOM 5.3.4.5.

You may encounter situations when management invites outside individuals to observe the inspectional process (e.g., representatives from the press, trade associations, congressional staff, other company officials). As discussed in Section 5.1.4.3 of the IOM, the presence of representatives invited by the firm should not disrupt the inspectional process. You are to continue the inspection in a reasonable manner.

If the firm allows invited individuals to photograph, videotape, or prepare audio recordings during the inspection, you should make every effort to protect privileged information in your possession. However, it is the Agency's position that it is the firm's responsibility to protect confidential and/or proprietary information observed or recorded by those individuals invited by the firm. Where applicable, refer to IOM 5.3.5 for additional procedures on how to prepare your own recording in parallel with the firm’s recording.

### 5.2.7.2 - Refusals of Requested Information

Should management refuse to provide any reasonable request for information, which is not specifically required by the law, determine the reasons for the denial and report the details in the EIR. Types of refusals of interest to FDA and refusal codes to be entered in FACTS are listed in the FDA Data Codes Manual. Refusal codes' data are used when reporting to Congress. See IOM 5.2.5.4 for instructions in dealing with hostile and/or uncooperative interviewees.

### 5.2.8 - CONSUMER COMPLAINTS

Prior to conducting any inspection, you should review the FACTS system and the factory jacket becoming familiar with all FDA Complaint/Injury forms. Be especially alert for ones marked "Follow-Up Next Inspection" and make sure you investigate these during your inspection.

During the inspection, discuss these complaints with management without revealing the complainant’s name(s). Determine if the firm has had similar complaints on the same product. Determine what action the firm has taken to identify the root cause of the problem and to prevent a recurrence in the future. See IOM 5.10.4.3.11 for reporting instructions.

### 5.2.9 - INTERVIEWING CONFIDENTIAL INFORMANTS

When you are faced with a situation involving sources of information who want to remain anonymous, please contact your supervisor and follow the procedures here. In addition, refer to IOM 5.2.1.2 regarding your personal safety. If your management concurs with the decision to utilize a confidential source, it is particularly important you take the necessary steps to keep the identity of the source, and any information which could lead to the identity, confidential. For purposes of this subchapter, a confidential source is a person who provides information that may be of assistance to FDA without necessarily becoming a party to the actual FDA investigation. If you believe the information provided by the source could lead to a criminal investigation, please contact the Office of Criminal Investigations (OCI).

### 5.2.9.1 - How to handle the first contact

When you interview a person who may become a confidential source use the following procedures:

1. **Type of meeting.** Try to schedule a personal interview with the person rather than a telephone interview. At a face-to-face interview you can assess the person's demeanor, body language, overall presentation, and truthfulness.

2. **Meeting location.** The place and time of the interview should be the choice of the person, unless there is a concern with personal safety. If the person’s suggested location is unsuitable, the investigator should suggest the location. When you conduct the interview off FDA premises, notify your supervisor of your destination, purpose, and estimated time of return. When an off-site interview has been completed, check-in with your supervisor.

### 5.2.9.1.1 - INTERVIEWING METHODS/TECHNIQUES

It is strongly recommended you have two investigators conduct interviews of a confidential source. The lead
involving the interview, while the second investigator takes notes and acts as a witness to the interview. You should:

1. Prepare carefully for the interview. The investigators should develop the questions they intend to ask the person during the interview, e.g., "establish motivation," and record and number the questions to be asked in their diaries prior to the interview. This preparation assists in documenting the interview process and reduces the amount of note taking needed during the interview. The investigators also should discuss their interviewing strategy, and determine the method by which they will consult with each other during the interview and (during extensive interviews) share the interviewing and note-taking responsibilities;

2. Have the person tell the story chronologically, placing complex situations into logical order; and

3. If the person makes allegations, ask him or her how he or she knows the allegations are true.
   a. How were they in a position to know?
   b. Did they personally see, hear, or write about the information/incident?
   c. Can they provide proof of the allegations?

5.2.9.1.2 - ESTABLISH MOTIVATION

At the end of the interview ask the person why he or she is divulging this information. This may reveal their motive(s):

1. Is the person a disgruntled current or former employee who harbors a grudge?
2. Is the person looking for some type of whistle-blower reward or notoriety?
3. Does the person just want to do the right thing?
4. Is the person involved in actual or prospective litigation about or related to the information?

5.2.9.1.3 - ANONYMITY

If the person is requesting anonymity, inform him or her FDA:

1. Will not divulge his or her identity, the occurrence of the interview, or the sensitive information provided to FDA if the information could lead to the identity of the person, unless FDA is required to disclose the information by law, e.g., the investigation leads to a hearing or trial and he or she is required to testify, and
2. Will try to corroborate all information provided by the person, minimizing the chances he or she must later testify. However, testifying remains a possibility.

Ask the person for names of other persons who might be willing to speak with you about the allegations and corroborate their story.

5.2.9.2 - Protect the Identity of the Source

Collection of information. Obtain sufficient personal information necessary to enable you to contact the person for follow up if needed. However, to maintain the confidentiality of the person, do not include the person's identifier information such as gender, name, address, and phone number in the memorandum of interview. You should assign the confidential source a code name or number and use the identifier in memoranda and other communications relating to the confidential source (see IOM 5.2.9.2.2 item 2).

5.2.9.2.1 - ACCESS

Know who is authorized by District procedure to access the information, and restrict access by others accordingly. Share the minimum amount of information necessary to meet the purpose of the disclosure.

5.2.9.2.2 - STORAGE REQUIREMENTS

Each district should establish procedures, in addition to those listed below, to properly store confidential information. The following list contains information related to storage procedures.

1. Use security measures necessary to protect the confidentiality of personal information, whether it is in hard copy or electronic form, on FDA premises, in an FDA home-based computer, or in any other form. Use whatever means necessary and appropriate to physically safeguard the information, such as storing in a safe, or locked file cabinets, or password-coded computers, etc.

2. When referring to the source in any manner (orally, in writing, electronically, etc.), consider using code to identify the source. For example, use a number rather than the individual's name, to identify the source. Personal privacy information should be safeguarded to the extent allowed by law. Use discreet subject headers in the file labels as appropriate.

3. Remove personal information from a file only after you have noted in the file your name, date, etc. Promptly return that information to the file.

5.2.9.2.3 - DISCLOSURE

Do not disclose information from or about the source, unless the disclosure complies with the law and FDA's procedures. Do not share non-public information outside of the Freedom of Information (FOI) process, unless the sharing is done according to our regulations and procedures. Refer FOI requests to your FOI officer (see item 3 below). See also IOM Subchapter 1.4. The following information relates to disclosures of information from or about a confidential source.

1. Make duplicates of the personal information only to the extent necessary for authorized disclosure (inside or outside of FDA). Do not leave the copy machine unattended.

2. Make only authorized disclosures of the information, regardless of the manner of disclosing (oral, written, etc.). Do not use mobile telephones or leave voice mails with the information. Avoid transmitting the non-public information by facsimile or e-mail.

3. If you receive a FOI request for information from or about a source consult with your supervisor immediately Disclosure to a non-FDA government official of information from or about a source may be disclosed only if permitted by law and FDA procedures,
and after consulting your supervisor and, if needed, OCC.
4. Immediately retrieve information from or about a source if inadvertently disclosed.

5.2.9.2.4 - DESTRUCTION

Destroy personal information by shredding or similar means which physically destroys the record and/or, if the information is in electronic form, makes it unreadable.

After a matter has been referred to the Office of Chief Counsel (OCC) for litigation or enforcement action, consult with OCC if you are interested in contacting the source.

5.2.10 - ROUTINE BIOSECURITY PROCEDURES FOR VISITS TO FACILITIES HOUSING OR TRANSPORTING DOMESTIC OR WILD ANIMALS

This section is FDA's guidance when you visit any type of facility where any domestic or wild animals are housed or transported. If a firm has more restrictive controls, follow those in addition to the controls cited below as long as they do not interfere with your assignment needs. The controls and procedures are intended to prevent you from becoming a vector or carrier of animal diseases, to prevent the spread of animal disease, and to set a good example for stockmen, growers and industry servicemen.

A number of chronic diseases, such as Johne's Disease, bovine virus diarrhea (BVD) and others exist in domestic animals which you can unknowingly spread. Any inspectional contact with herds of livestock (including poultry) or non-domesticated animals exposes you to potential claims of introducing or spreading disease. This could occur between sections of a single site, such as poultry houses, or between different sites or farms. The potential also exists for the introduction of disease from an animal processing plant, such as a slaughterhouse or renderer to a live animal facility. You can prevent this by following appropriate cleaning and disinfection steps between facilities. Generally, a break of 5 days or more between sites is sufficient to eliminate concern about transmission of infectious agents.

These precautions, biosecurity measures, are necessary in two types of situations. The first is when there is no known disease present and your actions are precautionary. This section primarily addresses those kinds of activities. The other situation involves known or suspected disease outbreaks or more notorious disease conditions such as salmonella in eggs, infectious Laryngotracheitis, foot and mouth disease, vesicular stomatitis, and blackhead which can be highly contagious and spread from one group of animals to another by movement of people and objects between infected and non-infected groups. In these cases, special precautions must be taken to make sure you are not an unknowing vector for the spread of disease. See IOM 5.2.10.3.

If you will only be inspecting an office or house away from areas where animals are housed or kept, clean and suitable street attire may be sufficient. Be aware if you visit any area of a facility where animals have been, you should always sanitize, clean or change footwear and it may be necessary to change outerwear before visiting another animal site to prevent any possibility of transmission of disease.

Your vehicle may also transport infection if you drive through contaminated areas and may require frequent cleaning between sites.

5.2.10.1 - Pre-Inspection Activities

When you know you are going to visit or inspect any animal production or holding facility, consider contacting the State Veterinarian and/or the Regional APHIS office to determine if there are any areas in the state under quarantine or special measures to control animal diseases. APHIS office locations can be found on their website. The State Veterinarian will be listed under Government Listings in your phone book and is listed at this website. Regional Milk Specialists frequently working with State counterparts in the Interstate Milk Shippers program should contact these sources at least quarterly for updates. Ask for any special controls or procedures they recommend. Follow any guidance they offer in addition to the precautions in this section. You should also consider pre-notification of the facility following guidance in IOM 5.2.1.1, Pre-Announcement, unless your assignment does not allow pre-notification. If you elect to pre-announce the inspection, in addition to the normal contact, ask to speak with the person at the facility responsible for their biosecurity measures and find out what they require of employees and visitors. If their requests do not interfere with your ability to do your job, follow their requests as we do when inspecting sterile manufacturing facilities.

Make sure your vehicle is clean and has been recently washed. Commercial car washes are adequate as long as you check to make sure any dirt, manure or other debris, which may be present from a previous site, has been removed. Some facilities may require additional disinfection of tires upon entry to the premises. Ensure tires and floor mats are clean. Consider designating places in your vehicle for storage of clean, unused supplies and dirty or used supplies.

In addition to your normal inspectional tools, obtain the following equipment and supplies from your district:
1. Laundered or disposable coveralls or smocks (coveralls are suggested because they give better coverage). If you are going to visit multiple facilities in one day or trip, obtain sufficient quantities so you can change into clean or unused clothing between each site.
2. Disposable plastic gloves, rubber boots, which can be sanitized, and disposable shoe/boot covers. Rubber boots over which you place disposable shoe/boot covers are preferred.
5. Reusable cloth or plastic laundry bag(s) for clothing to be laundered. (Disposable bags can be used.)
4. Soap, water and disposable or freshly laundered individual hand (or paper) towels.
5. Sanitizing solution(s) and equipment (brushes, bucket, tray, measuring devices, etc.) to permit you to properly sanitizing hands, boots, equipment and your vehicle. Most disinfectants will require removing organic matter before use and good brushes are essential to remove dirt from boots and other objects.

Make sure any equipment you take with you has been thoroughly cleaned and sanitized as necessary. Clip boards, briefcases, flashlights, inspectional sampling tools, coolers, brushes, buckets and other objects should be cleaned between uses as necessary and between visits to any suspected infected facilities. Disposable equipment should be used to the fullest extent possible.

Maintain copies of any applicable Material Safety Data Sheets (MSDS) for disinfectants with you in your vehicle. If the firm's management requests information on the disinfectants you are using, they may read or copy these MSDS. Be familiar with the instructions and precautions concerning use of disinfectants. Any disinfectant should be effective against known or suspected microbiological agents.

In the event of a foreign animal disease, contact the USDA, APHIS Veterinary Services area Veterinarian in Charge for additional precautions and procedures to follow. (See 5.2.10.3)

5.2.10.2 - General Inspection Procedures

Always begin each day with a clean vehicle free from any visible dirt or debris. During the day, take precautions to minimize contamination of your vehicle. If your vehicle becomes obviously dirty with adhering mud or manure, clean it before visiting another animal facility. When you arrive at a facility where animals are located, check to see if there are designated parking spots or pads for visitors. If so, park your vehicle there unless directed otherwise by the firm. If there is no guidance, park well away from all areas housing animals. When you arrive, inquire about or reconfirm any biosecurity measures the firm employs. Confirm your actions are suitable and follow expectations of the facility when this does not interfere with your inspection ability. Follow steps requested by the firm to remove contamination from vehicles, which may include troughs or pools of disinfectants for tires or other control measures. Avoid driving through manure, mud or wastewater at these sites.

In general, entry to animal housing or feeding areas, corrals, calf pens, hospital pens or special treatment facilities should be avoided unless the assignment requires their inspection or there are specific reasons requiring entry. If you must visit the feeding area occupied by livestock or birds, first determine if any groups are infected with disease. Arrange to visit the known non-disease areas first. Do not handle any animals unless official duty requires such contact. Before leaving the area where you parked your car, put on protective clothing as described and proceed with the purpose of your visit; sanitizing hands (and gloves if worn) and boots as necessary during the visit or inspection.

General procedures:
1. Wear rubber boots or other suitable footwear, which you disinfect upon arriving at the site and prior to departure. It is preferable to also place disposable foot coverings over your footwear, regardless of the type, after you have disinfected them. If the firm has footbaths, use them. Boots and footwear should be disinfected with any of the agents identified at the end of this subsection using a good brush. Clean and disinfect the brush(es) and bucket you use for these activities.
2. Wash your hands with soap and water. If you are visiting a facility where a known animal disease is present or the firm's biosecurity protocol requires, wear disposable gloves.
3. Wear disposable or freshly laundered coveralls, when appropriate. Some facilities may provide disposable coveralls and require visitors to shower in and shower out at their facilities. If requested by the firm and facilities are provided, you should follow those requests.
4. Wear appropriate head coverings, as necessary. If you wear a head covering, clean and disinfect between facilities or use disposable head coverings.
5. Minimize any materials you carry with you such as notebooks, flashlights, etc. to what is required. Consider keeping these things in clean plastic bags or containers between uses. Disinfect any of these types of items as best you can between visits to facilities or between different animal-housing areas.
6. If you are visiting production units with animals of multiple ages, always try to work from the youngest to the oldest.
7. Avoid direct contact with livestock or wild animals, bodily fluids or animal byproducts when visiting facilities.
8. Regional Milk Specialists, Milk Safety Branch and State Training Team staff frequently working with State counterparts in the Interstate Milk Shippers program shall follow any biosecurity measures the firm employs, any biosecurity measures the State employs, and as a minimum shall follow the coded memoranda issued by CFSAN Milk Safety Branch on this subject.

Upon completing your assignment in a given animal area, return to the same area where you donned protective clothing. Remove disposable shoe/boot covers and gloves, if applicable, and place them in a disposable paper or plastic bag. Clean and sanitize boots/footwear. Remove the protective clothing, if applicable, by peeling it off inside out. (This keeps the surfaces exposed to contamination on the inside.) Unless the firm's biosecurity plan prohibits removal of waste from their premises, all waste should be disposed of by the investigator as follows: Place all disposable items in a disposable, nonporous bag for appropriate disposal according to State and/or local
5.2.10.3 - Special Situation Precautions

If you are required to inspect or visit a facility known or suspected to be involved in a contagious animal disease outbreak or otherwise identified as having diseased animals, contact the Center for Veterinary Medicine and/or Center for Food Safety and Applied Nutrition for additional precautions which may be necessary before you visits these sites. Your activities may be limited to visiting a single site in a day, taking extra-ordinary decontamination steps, ensuring you do not visit or inspect another facility for 5 or more days following the visit to the contaminated site or other steps. APHIS may have special restrictions or precautions for you to follow. The State Veterinarian may also request you follow additional requirements. During inspections of poultry operations where salmonella contamination is known or suspected, you should make sure you contact CFSAN directly for specific procedures to follow. Additional decontamination steps will be required.

5.3.1 - TECHNIQUES

The recognition, collection, and effective presentation of admissible evidence is essential to successful litigation. Tangible evidence is required to support your observations and reports of violative conditions.

Although the inspectional procedures to detect adulteration and contamination, etc., are described under specific headings in the IOM, the same procedures and/or techniques may also apply to other areas. For instance, the procedures to detect contamination from filth, insects, rodents, birds, etc., described in IOM section 5.4.7 may also apply to drugs or other products. Your experience and training assists you in making this transition and enables you to detect possible violative conditions.

Keep in mind the policy annunciated in the 4/23/1991 memorandum from the Director, Office of Compliance: The lack of a violative physical sample is not a bar to pursuing regulatory and/or administrative action providing the CGMP deficiencies have been well documented. Likewise, physical samples found to be in compliance are not a bar to pursuing action under CGMP charges.

5.3.2 - FACTORY SAMPLES

Samples of raw materials or finished products collected during inspections provide the necessary key to establish routes of contamination. They also document the character of products packed prior to the inspection. Collect Factory Samples for laboratory examination only when they contribute to confirming the suspected violation. Be selective since negative reports of analysis of food samples are required under Section 704(d) of the FD&C Act [21 U.S.C. 374 (d)] to be furnished to the firm and might give management a false picture of the firm's operation.

When possible collect duplicate subsamples to provide for the 702(b) portion of the sample. See IOM 4.3.2.1 and 4.3.7.4.1 for additional guidance and 21 CFR 2.10 for exemptions regarding the collection of duplicate portions.

5.3.3 - EXHIBITS

Impressive exhibits are extremely effective and important forms of evidence to establish existence of violative conditions or products. They should relate to insanitary conditions contributing or likely to contribute, filth to the finished product, or to practices likely to render the product injurious or otherwise violative. Diagrams of the establishment, floor plans, flow charts, and schematics are useful in preparing a clear concise report and in later presentation of testimony. A small compass is useful in describing exact locations of objectionable conditions in the plant, in your diagrams, and locations from which samples were taken, etc.

SUBCHAPTER 5.3 - EVIDENCE DEVELOPMENT

Describe and submit under one INV Sample Number all exhibits (except photographs) collected during the
inspection or investigation. Identify and number individual subs and officially seal all samples collected.

Examples of exhibits include:
1. Live and dead insects.
2. Insect frass, webbing, and insect chewed materials; nesting material of rodents and/or other animals; and other behavioral evidence of the presence of insects, rodents and other animals.
3. Samples of components or ingredients, in-process materials and finished products or dosage forms.
4. Manufacturing and control devices or aids.
5. Physical samples if possible and practical or, photographs with descriptions of scoops, stop-gap expediencies, other unorthodox manufacturing equipment or makeshift procedures. If photos are taken, follow the procedures described in IOM 5.3.4.
6. Evidence showing the presence of prohibited pesticide residues. A method of swabbing for prohibited pesticide residues was published in Laboratory Information Bulletin # 1622. Excerpts are quoted as follows:

**5.3.4 - PHOTOGRAPHS**

Photos taken during EI’s are not classified as INV Samples. They are exhibits. No C/R is used for photos taken unless the photos are part of an Official Sample. See IOM 4.1.4 for information on Official Samples.

Since photographs are one of the most effective and useful forms of evidence, every photo should be taken with a purpose. Photographs should be related to insanitary conditions contributing or likely to contribute filth to the finished product, or to practices likely to render it injurious or otherwise violative.

CAUTION: Evaluate the area where flash photography is contemplated. Do not use flash where there is a potentially explosive condition; e.g. very dusty areas or possible presence of explosive or flammable vapors. In these situations use extremely fast film and/or long exposure time instead of flash.

Examples of conditions or practices effectively documented by photographs include:
1. Evidence of rodents or insect infestation and faulty construction or maintenance, which contributes to these conditions.
2. Routes of, as well as, actual contamination of raw materials or finished products.
3. Condition of raw materials or finished products.
4. Employee practices contributing to contamination or to violative conditions.
5. Manufacturing processes.
6. Manufacturing and various control records showing errors, substitutions, penciled changes in procedure, faulty practices, deviations from GMP’s, NDA’s, or other protocols, altered or inadequate assays or other control procedures and any variation from stated procedure. See IOM 5.3.8.2 for identification of records.
7. Effluent contamination of water systems. See IOM 5.4.3 for techniques in photographing this type of contamination.

When photographing labels, make sure your picture will result in a legible label with printing large enough to be read by an unaided eye. Photograph whitewashed documents by holding a flashlight against the whitened outer side and taking a close up photo of the reverse using high-speed film. This will produce a photo with a mirror image of the whitened outer side.

If you use a Polaroid camera or color slide film, explain the facts in your EIR or on the C/R to alert reviewers that there are no negatives.

5.3.4.1 - In-Firm Photographs

Do not request permission from management to take photographs during an inspection. Take your camera into the firm and use it as necessary just as you use other inspectional equipment.

If management objects to taking photographs, explain that photos are an integral part of an inspection and present an accurate picture of firm conditions. If management of a drug firm does not give a reasonable explanation for its objection, such as a showing that the chemical properties of products manufactured at the facility are such that taking photographs would adversely affect product quality, you may advise management that the refusal may constitute a limiting of the inspection under Section 501(j) [21 U.S.C. 551(j)] of the FD&C.
Advise management the U. S. Courts have held that photographs may lawfully be taken as part of an inspection.

If management continues to refuse, provide them with the following references:

   This Supreme Court Decision dealt with aerial photographs by EPA, but the Court's language seems to address the right to take photographs by any regulatory agency. The decision reads in part, **"When Congress invests an agency with enforcement and investigatory authority, it is not necessary to identify explicitly each and every technique that may be used in the course of executing the statutory mission."**


If management refuses, obtain name and contact information for the firm's legal counsel, and advise your district management immediately. If the firm does not have legal counsel on retainer, collect the name and contact information for the most responsible individual. District management will inform their ORA Regional Counselor in the Office of Chief Counsel (OCC) of the situation, and OCC will then contact the firm's legal counsel or most responsible individual to discuss FDA's legal right to take pictures during inspections. OCC will relay the results of this conversation to district management. If you have already taken some photos do not surrender film to management. Advise the firm it can obtain copies of the photos under the Freedom of Information Act. See IOM 5.3.4.5.

### 5.3.4.2 - Photo Identification and Submission

One of the most critical aspects about photographs or videotapes is the ability for the agency to provide testimony clearly verifying the authenticity of the conditions depicted in the photograph or video. It makes no difference if the photo is a 35 mm print from acetate negatives, a Polaroid photo, a digital photo or video taken with a video recorder. You must create a trail, starting with the taking of the photo, confirming its original accuracy and establishing a record describing the chain of custody. To do this, you must make sure each photograph is described in your regulatory notes in sufficient detail to assure positive correlation of the photo or video with your inspection findings. One way you can do this is to photograph a card with your name, district address and phone number as the first frame or picture on a roll of film or in the digital record. This will help identify the film or file and assist in tracking if it is lost or becomes separated from its identification envelope during processing or storage. Proper procedures will also allow the agency to provide evidence confirming the authenticity of the photographs or video recording in the event you are not able to testify personally.

#### 5.3.4.2.1 – FILM BASED PRINTS

Identify each print on the margin with exhibit number, firm name (or DOC Sample Nos., if DOC Sample), date taken or inclusive dates of inspection, and your initials. Do not place any identifying marks on the picture area of the print. (Some photo developing firms are supplying borderless prints. For this type print, place identification along the back bottom edge of the print and mount the print so the identification can be read without removing the print from the mounting paper. Place a narrative description on the mounting paper next to the print and attach as exhibits to the EIR and/or route with other records associated with a DOC Sample.)

#### 5.3.4.2.2 - COLOR SLIDE IDENTIFICATION

If color slides are used, identify each slide, in the same manner as for prints. Districts may have special mounting frames for color slides, so the narrative description of each slide must be in the body of the report with proper reference to exhibits, or, each description may be placed on sheets of paper following the mounting frames and properly referenced.

#### 5.3.4.2.3 - NEGATIVE IDENTIFICATION

Identify the edge of at least two negative strips, with the same information as for prints using a 3/16" strip of pressure sensitive tape. Place all negatives in a FDA-525 envelope. Complete blocks 2, 3, (4 if DOC Sample), 5, 7, and 12 and seal with an Official Seal, FDA-415a. If negatives are not part of a DOC Sample, enter firm name in the Sample Number block.

As applicable, submit the sealed FDA-525 or envelope as an exhibit to the EIR, with the Investigative Report as an attachment, or with the other associated records/documents with a DOC Sample.

#### 5.3.4.2.4 - VIDEO RECORDINGS

Handle and protect the original video record just as if it were a photograph negative. Unused videotapes should generally be used to capture the video and, for subsequent copies of the original recording. Write-protect and identify the original videotape with a label with the firm name (or Sample number if it is being submitted as part of an official sample), date taken, and your initials. Officially seal the original videotape in a FDA-525 envelope or similar envelope. If you use a larger, unfranked envelope, identify the envelope with your name, title, home district, date, firm name, firm address (include zip code), description of the contents of the envelope, and marked in large, bolded letters "STORE AWAY AND PROTECT FROM MAGNETIC FIELDS." You may place more than one videotape in a single FDA-525 as long as you state on the envelope how many videotapes are in the envelope. If the original envelope is opened, document the chain of custody and use new seal(s) after each entry to the envelope.
If you perform any editing of the recording, you should only perform this on a copy of the original video recording to prevent possible damage to the original. Document in your regulatory notes you made a copy of the original and verified the copy is an accurate copy of the original video you took. This “original copy” should be treated just as if it is the original. When you sign the report, memorandum or other agency document, your signature certifies you are saying the content of the document, including any video recordings, is true and accurate to the best of your ability.

As applicable, submit the officially sealed FDA-525 or envelope as an exhibit to the EIR, with the Investigative Report as an attachment, or with the other associated records/documents with a DOC Sample.

5.3.4.2.5 – DIGITAL PHOTOGRAPHS OR VIDEO RECORDINGS

Prior to the year 2000, FDA investigators traditionally worked with silver acetate photographic film or used analog video tapes. Early digital cameras recorded photographic images directly to floppy disks or mini-CDs in which the evidence could be handled like photographic negatives.

The important difference today is digital cameras are capable of recording high resolution images on the order of twenty to thirty megapixels. The corresponding image file sizes can be over fifteen megabytes when using uncompressed file formats. To cope with the increased file sizes, digital camera manufacturers have introduced non-volatile flash memory cards which can record digital images, delete images, and be recorded over and over again. This presents a new issue since the original digital images, which are captured at the moment when the images are recorded on the memory card, will be copied at a later time to a CD-R or other permanent storage media. Due to the cost of flash memory cards and the large file sizes, it is not feasible to purchase new memory cards for each inspection/investigation as you did using photographic film. You will be working with an “original copy” of the images which have to be copied in the exact format to a CD-R or DVD-R as they were originally recorded on the flash memory card to preserve the chain of custody. The term “other permanent storage media” includes the hard-drive on the work computer/laptop of the investigator, and not a shared or personal computer. In order to preserve the chain of custody, it is acceptable to transfer the images from the flash memory card onto the hard-drive and then burn the images onto a CD-R or DVD-R, so long as the images have not been altered in any way before being burned onto the CD-R or DVD-R.

In the same manner, digital video recordings may involve the use of different media types such as tapes, CD-Rs or DVD-Rs, or built-in hard drives. If you cannot handle the original video recording as in IOM 5.3.4.2.4, you will need to create an “original copy” of the video recording.

Despite the differences in photographic film and digital technology, you are responsible for collection, handling, documenting the chain of custody, storage, and submission of your evidence in a manner where you can testify to its authenticity in a court of law. See IOM 5.3.4.2 and 5.3.4.3.

5.3.4.2.6 – GLOSSARY OF DIGITAL TERMINOLOGY

Some basic terminology is used when referring to digital devices in IOM 5.3.4.2.4, 5.3.4.2.5, 5.3.4.3.

5.3.4.2.6.1 – Digital Data

Electronic data in binary form consisting in its simplest form as “1”s and “0”s. A computer interprets data by whether the state is on (“1”) or off (“0”).

5.3.4.2.6.2 – Analog Data

Information captured in a directly measurable signal versus an analog signal converted and stored in binary.

5.3.4.2.6.3 – Memory Card

Any non-volatile memory media that can be removed and which retains data without the need for electrical power. Examples of current memory cards are: Compact Flash (CF), Secure Digital (SD), Memory Stick (Sony), and Extreme Digital (xD).

5.3.4.2.6.4 - Original

The file recorded by a digital device on digital storage media at the moment in time when the user takes a picture or makes a recording. This concept is similar to a film camera where the photographic film records the image when exposed by light. The film image negatives produced when the film is developed are considered the originals and prints are considered copies. See IOM 5.3.4.2.1 and 5.3.4.2.3.

5.3.4.2.6.5 – Original Copy

An exact copy of the original file recorded by the digital device (camera, video recorder, etc.). The original copy will retain all the characteristics of the original and is indistinguishable from the original.

5.3.4.2.6.6 - Permanent Storage Media

A media format in which the digital files cannot be altered once written. Examples are CD-Rs, DVD-Rs and other approved media.

5.3.4.2.6.7 - Time/Date Stamp

The internal clock within the camera which records the time/date information on the image file. Set the time/date stamp for the location where the photographs or videos are being taken. In this usage, the time/date stamp does not refer to imprinting the time/date stamp within the photographic image although the time/date stamp can also be imprinted on the photograph as some film cameras could do.
5.3.4.2.6.8 - Working Copy

A copy of the original copy used when you need to make additional copies for your report, sample C/R. Creating a working copy decreases the chance the original copy is damaged.

5.3.4.3 - Preparing and Maintaining Digital Photographs as Regulatory Evidence

Assure and protect a digital photo's chain of custody (and authenticity) following this procedure:

1. Prior to using the digital camera, verify the date and time stamp is correct and there are no images stored on the memory card. Reformat the memory card using your camera’s reformat command to delete any images not related to your current assignment. Depending on your inspection/investigation, camera, and memory card capacity you should consider bringing more than one memory card if possible.

2. Handle your camera and the memory cards in a manner to protect your evidence and maintain the trail of the "chain of custody" for the evidence you have collected. For example, keep the camera and memory cards in your personal possession at all times or held under lock/key in a secure storage area. Also, keep any additional memory cards containing images in your personal possession until transferred to permanent storage media. Where necessary, document these facts in your regulatory notes or written report (EIR, CR etc).

3. As soon as practical, create an original copy of the digital photos. Some older FDA cameras will capture images directly to a (Write-once Compact Disk Recordable (CD-R)); in this case, the CD-R from these cameras becomes the original CD-R. Identify, date and initial the CD-R as an original image record. If a CD-R/W was used, copy the images to a CD-R to create an original copy with files that cannot be altered. Follow additional instructions for creating and finishing a CD-R in step 4 below.

4. If the camera requires downloading of images to a CD-R or other media, download all the images from the digital camera to an unused CD-R or other electronic storage media to create an original copy. If there was more than one memory card used, use a separate CD-R for each memory card. The storage capacity of a CD-R is about 650 MB; thus, more than one CD-R may be needed to create an original copy of your memory card depending on your camera’s resolution, the storage capacity of your memory card, and the number of pictures taken. The images should be transferred in a file format maintaining the image resolution at the time the image was captured. If possible, avoid the use of any file compression in transferring the images to the CD-R. Prior to preparing the CD-R or transferring image files, verify that the computer you are using is set to the correct date and time. Make the CD-R permanent in a format readable by any CD-R reader. Prior to making the working copy from the original copy, identify the original copy with the same information as in IOM 5.3.4.2.1. It is important to identify the original copy as soon as possible to prevent possible mix up of the original copy with any working copies.

5. Use a permanent CD safe marker to identify the original copy CD-R. Do not use ball point pens or similar tipped markers since the CD-R may be damaged. See the NIST document, “Care and Handling of CDs and DVDs - A Guide for Librarians and Archivists”. Figure 12, page 23 shows where to identify the CD-R.

6. Where applicable, document in your regulatory notes the verification and identification of each photographic image comparing them to your regulatory notes, which were recorded at the time the photographs were taken.

7. Make only one working copy from each original copy and make any additional working copies using the initial working copy. No more than one copy should be made from the original copy in order to preserve the original copy a pristine set. After making the initial working copy, place the original copy in a suitable package, officially seal and store the original copy (CD-R or other electronic storage media) until submitted with the written report (EIR, CR etc). If the images are captured or transferred to disc(s), refer to IOM 5.3.8.3 for the handling of disc(s). If possible, the investigator (who took the photos and will authenticate them at trial) should store the sealed CD-R or other electronic storage media until it is submitted with the written report. If you break the seal for any reason, document this on the broken seal, in your regulatory notes or written report, and reseal the package with a new official seal. See IOM 4.5.4.5.

8. Working copies should be used to print photos, insertion into an EIR, cropped, otherwise edited or to be included in a referral.

9. Document in your regulatory notes or written report (EIR, CR, etc.) any steps taken for any unusual editing of original photo images. For example: Superimposing over an important area of the image, image enhancement, composite images, etc.

5.3.4.4 - Preparing Digital Photos for Insertion in a Turbo Establishment Inspection Report (EIR)

Digital photos taken during an inspection can be inserted into the body of a report in Turbo EIR or can be printed and attached to the EIR as an exhibit. Inserting digital photos can dramatically increase the file size of the Turbo EIR document. To maintain a minimum Turbo EIR document file size, the following is recommended: Do not open a digital picture/photo and use copy and paste to insert the picture/photo into the Turbo EIR document. Instead, save pictures/photos in a JPEG image format (.jpg file name extension) in a separate folder in preparation for inserting into Turbo EIR. Then resize all the JPEG pictures to a reasonable image file size. To do this,
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1. Open the folder with all the pictures that may be inserted into the Turbo EIR document.
2. Hold the control key down and left click to select each image file to be resized.
3. Right click, choose resize pictures. See Exhibit 5-6.
4. Select a size -- click on Small (fits a 640 x 480 screen), and click OK. Selecting one of the other screen sizes will also work with the exception of "Handheld PC (fits a 240 x 320 screen)"
5. New resized files will be created within the same folder. Each original file will be maintained. Each new resized file will be renamed as original file name (Small).jpg to differentiate it from the original file.
6. The resized pictures/photos are now ready for insertion into the Turbo EIR document. Remember to maintain the original image files and not the resized digital image files for submission with your report, forms, and exhibits to the official establishment file.

To insert a picture into the Turbo EIR document:

1. Open the Turbo EIR document Position cursor to where you want to insert the picture.
2. From the menu bar, click on Insert, choose Picture, click on From File, find and select folder with resized pictures to be inserted. See Exhibit 5-7.
3. Double click on the resized picture to be inserted.
4. Picture inserted into the Turbo EIR document can be made larger or smaller by clicking on the picture and grabbing the corner of the picture frame and dragging to achieve the desired size.

Alternative method: Digital photographs can also be submitted as Exhibits to the EIR. A narrative description may be placed below the digital photograph. Include the photo number, the date photo was taken and by whom, and a brief description of what the photo depicts. See also 5.3.8.2 - Identification of Records Collected.

Photographs can be resized using Microsoft Office Picture Manager. See Exhibit 5-8 which shows the "resize" menu option.

NOTE: When any digital photos are used in an EIR, submit the original or original copy of the camera images following procedures as outlined in IOM 5.3.4.3 – Preparing and Maintaining Digital Photographs as Regulatory Evidence.

5.3.4.5 - Photograph Requests

Do not routinely advise firms they may have copies of photos. However, if management of the firm initiates the request, advise them it is possible to obtain copies of photographs taken in their plant under the Freedom of Information Act. Any request should be sent to The Food and Drug Administration, at the address listed on the FDA 482 or FDA 483. The firm must bear the cost of duplicating the photographs.

Since photographs are records in an investigative file, they are not available under the Freedom of Information Act until the file is closed.

Do not discourage firms from taking their own photographs at the same time and of the same scenes as you.

5.3.5 - RECORDINGS

Under normal circumstances recording devices will not be used while conducting inspections and investigations. However, some firms are now recording and/or videotaping, the inspection and/or the discussion with management portion of the inspection. These firms should be advised we do not object to this procedure, but we will also record the discussion to assure the accuracy of our records. Occasionally a firm’s management may record the serving of an inspection warrant or, in a hostile situation, may want to record everything. In such cases, depending on the circumstances, you may prepare your own recording in parallel with the firm’s recording. Do not depend on the firm to provide a duplicate of their recordings.

Use a clear tape cassette and identify the tape verbally as follows:

"This is Investigator ____________ of the U.S. Food and Drug Administration speaking in the (state location) of (firm name) (dates). It is now a.m./p.m. on (date). Present are (list individuals present with title). This discussion is being recorded by both the representative of (firm name) and by me. We are going to discuss the inspectional findings of an inspection conducted at this firm on (inclusive dates)."

At the close of the discussion and prior to leaving the firm, the recording will be verbally identified as follows:

"This is Investigator ____________ speaking. It is now _______ a.m./p.m. on (date). This was a recording of the discussion with management at the conclusion of an inspection of (firm name and address) conducted on (dates)."

If the recording covers a different situation, the identification should be modified accordingly. If the representative of the firm refuses permission to record the discussion, continue with your discussion and report the facts in your EIR.

The tape cassette must be identified with the firm name, date of the inspection, and investigator’s name. Districts have the option of transcribing the tape and making the transcription an exhibit for the EIR. However, the tape itself must be made a permanent part of the EIR as an exhibit.

5.3.6 - RESPONSIBLE INDIVIDUALS

The identification of those responsible for violations is a critical part of the inspection, and as important as determining and documenting the violations themselves. Responsibility must be determined to identify those
persons to hold accountable for violations, and with whom the agency must deal to seek lasting corrections.

Document and fully report individual responsibility whenever;
1. It is required by the assignment,
2. Inspectional findings suggest the possibility of regulatory action, or
3. Background information suggests the possibility of regulatory action.

Under the Medical Device Quality System regulation (21 CFR 820.20), if the management at the firm is not exercising the controls required by the regulation, the deviations may be cited on your FDA 483.

5.3.6.1 - Discussion on Duty, Power, Responsibility

Duty - An obligation required by one's position; a moral or legal obligation.

Power - Possession of the right or ability to wield force or influence to produce an effect.

Responsibility - An individual who has the duty and power to act is a responsible person.

Three key points to consider are:
1. Who had the duty and power to detect the violation?
2. Who had the duty and power to prevent the violation?
3. Who had the duty and power to correct the violation?

5.3.6.2 - Inspection Techniques How to Document Responsibility

Always determine and report the full legal name and title of persons interviewed, who supplied relevant facts and the name/title/address of top management officials to whom FDA correspondence should be directed.

Obtain the correct name and correct title of all corporate officers or company officials. Obtain pertinent educational and experience backgrounds, and the duties and powers of the officers and employees in key managerial, production, control, and sanitation positions. Ascertain the experience and training of supervisory personnel, in terms that will describe their qualifications to carry out their responsibilities.

There are numerous ways to establish and document responsibility. Evidence may be obtained during interviews and record review specifically intended to determine responsibility. Cover and report items such as:
1. Organizational charts,
2. Statements by individuals admitting their responsibility or attributing responsibility to others,
3. Company publications, letters, memos and instructions to employees, and
4. The presence or absence of individuals in specific areas at specific, significant times, and their observed activities directing, approving, etc.

In order to establish relationships between violative conditions and responsible individuals, the following types of information, would be useful:
1. Who knew of conditions?
2. Who should have known of the conditions because of their specific or overall duties and positions?
3. Who had the duty and power to prevent or detect the conditions, or to see they were prevented or detected?
4. Who had the duty and power to correct the conditions, or to see they were corrected? What was done after person(s) learned of the conditions? Upon whose authority and instructions (be specific)?
5. What orders were issued (When, by whom, to whom, on whose authority and instructions)?
6. What follow-up was done to see if orders were carried out (when; by whom; on whose authority and instructions)?
7. Who decided corrections were or were not complete and satisfactory?
8. What funding, new equipment, new procedures were requested, authorized or denied in relation to the conditions; who made the requests, authorizations, or denials.

Duties and power related to general operations should be established to supplement the specific relationships to violations. Examples of operational decisions that indicate responsibility are:
1. What processing equipment to buy.
2. What raw materials to purchase.
3. What products to produce and what procedures to follow in production?
4. Production schedules - how much to produce, what to make, when to stop or alter production?
5. What production controls to be used?
6. What standards are set for products, raw materials, processes?
7. How to correct or prevent adverse conditions; how much to spend and whom to hire to correct or prevent adverse conditions; when to clean up?
8. How products will be labeled; what products to ship; label approval?
9. When to reject raw materials or products; when to initiate a recall; acceptable quality levels for products?
10. When to hire or fire personnel?
11. Who will accept FDA 482, Notice of Inspection; refuses inspection; accept Inspectional Observations, FDA 483?
12. Who designed and implemented the quality assurance plan; who receives reports of Q.A.; who acts or should act upon the reports?
13. Who is responsible for auditing other facilities, contractors, vendors, GLP sites, etc.?
14. In the firm's business relationships, who signs major contracts, purchase orders, etc.?
In some circumstances, documenting of individual responsibility requires investigative techniques that lead to sources outside the firm. These sources may include contractors, consultants, pest control or sanitation services, local health officials and others. Copies of documents between the firm and outside parties may help establish responsibilities. Do not overlook state officials as another possible source of information in selected cases.

During the course of the inspection you may observe persons who hold responsible positions and/or influence in the firm whose abilities or judgment may be affected by an obvious infirmity, handicap, or disability. If it is obvious the infirmity adversely affects the person's responsibilities or duties that are under FDA oversight, describe in your EIR the extent of the infirmity and how it relates to the purported problem or adverse condition.

5.3.7 - GUARANTEES AND LABELING AGREEMENTS

Review the Code of Federal Regulations, 21 CFR 7.12, 7.13, 501.100(d), 201.150, and 701.9, for information concerning guarantees and labeling agreements.

5.3.7.1 - Guaranty

Certain exemptions from the criminal provisions of the FD&C Act are provided where a valid guarantee exists as specified in Section 303(c) of the FD&C Act [21 U.S.C. 333 (c)]. Obtain a copy of any Food and Drug guarantee, which the firm claims to use relating to a violation noted during your inspection. No person may rely upon any guaranty unless he has acted merely as a conduit through which the merchandise reached the consumer.

5.3.7.2 - Labeling Agreement

Products regulated by FDA are normally expected to be completely labeled when introduced into or while in interstate commerce. Under certain conditions exemptions are allowed when such articles are, in accordance with trade practices, to be processed, labeled, or repacked in substantial quantity at an establishment other than where originally processed or packed. Sections 405, 503(a) and 603 of the FD&C Act [21 U.S.C. 345, 353(a), and 363] also provide exemptions from complete labeling for products.

5.3.7.3 - Exemption Requirements

To enjoy this exemption, the shipment must meet one of the following:
1. The shipper must operate the establishment where the article is to be processed, labeled or repacked; or
2. If the shipper is not the operator of the establishment, he must first obtain from the owner a written agreement signed by and containing the post office addresses of such persons and such operator and containing such specifications for the processing, labeling or repacking of such articles as will insure that such article will not be adulterated or misbranded within the meaning of the Act, upon completion of the processing, labeling or repacking.

Submit copies and dates of labeling agreements where unlabeled articles are shipped in interstate commerce.

5.3.8 - RECORDS OBTAINED

Many types of inspections and investigations require collection of copies of records to document evidence of deviations. In some cases, this may involve voluminous copies of Good Manufacturing Practice (GMP) records, commitments made in the Pre-Approval process, adherence to the requirements of the Low Acid Canned Food regulations or other areas. Copies of records are also obtained to document interstate commerce, product labeling and promotion, and to identify the party or parties responsible for a variety of actions. Copies of records can be obtained in paper format or electronic format. All records become part of the government's case should it go to litigation.

Normally, during litigation proceedings, the best evidence rule prevails in court, whereby the copy of the record in the custody of the government can be authenticated, if the original record is not produced by the custodian of the record.

It is imperative the government witness [usually the collector of the record(s)] be able to testify where, when and from whom the copies were obtained, and that the copy is a true copy of the source record, based on their review of the source record.

5.3.8.1 - Verification of Source Records

You must verify the copy of the record(s) you received is an accurate representation of the original or source record(s) so you are able to testify your copy is an exact duplicate of the original or source record. Record in your regulatory notes you authenticated copies of records and when, where, and from whom copies were obtained.

5.3.8.2 - Identification of Records Collected

Articles used as evidence in court cases must be identified so you can later testify the records entered as evidence are the very ones you obtained. This includes all records as noted in IOM 5.3.8, and any others for evidence in administrative or judiciary proceedings. When identifying and filing records, you must ensure the record is complete and no identification method or filing mechanism covers, defaces or obliterates any data on the record.

You must identify records submitted in support of an inspection or investigation, including records provided in an Establishment Inspection Report (EIR) or narrative memorandum. The identification must positively identify the specific copies you received during your inspection or investigation and to avoid any filing mix-up. If labels are
used to identify records, they must be permanently applied so any removal will be obvious.

You should identify records submitted with an EIR using the FEI number, firm name, , first and last date of the inspection, your initials exhibit number, and page numbers. When you collect a sample, each page of the copied records will become part of the collection report and should be identified as noted in IOM 4.4.5. Examples include records of interstate commerce, manufacturing deviations, label and labeling violations. Records submitted with a memorandum (see Exhibit 5-17) will include a phrase or firm or subject name to tie them to the investigation, the date(s) of the investigation and your initials. There are occasions when a single record may include hundreds of sheets of bound paper. Abbreviated methods of identification may be used for bound records by fully identifying the first and last few pages. In some cases, firm's clearly mark each page with the sequential and total pages number (e.g., page 6 of 10, 7 of 10, etc.) and this allows you to fully mark only a few pages in the beginning and end of the exhibit.

All pages must be identifiable if not in bound records. One example of a shortened method of identifying individual exhibits containing a large number of pages (usually more than 25) is to fully identify the first few and last few pages with at least the exhibit number, date and your initials. Then identify the remaining pages with the page number of the total page numbers, and your initials, e.g., "5 of 95 SHR". This may not be acceptable if you have more than one exhibit consisting of exactly 95 pages.

Whatever method is used, you must assure the record is complete and is always identifiable. This is so you can testify as to the "where", "when" and "from whom" the copies were obtained, and that the copy is a true copy of the source record based on your review of the source record. The identification method should allow any reviewer to determine if the record is complete or pages or parts are missing.

### 5.3.8.3 - Filmed or Electronic Records

When attempting to obtain records, you may find they are stored on microfilm, microfiche, or some form of a computerized management information system as electronic records. Paper or electronic copies of records obtained during the course of the inspection from these sources are handled the same as any records following procedures outlined in IOM 5.3.8, 5.3.8.1, and 5.3.8.2.

#### 5.3.8.3.1 - DEFINITIONS

Sections 5.3.8.3.1.1 to 5.3.8.3.1.5 contain definitions of some basic terminology used when handling electronic records as identified in section 5.3.8.

### 5.3.8.3.1.1 ELECTRONIC RECORDS

Electronic records are defined in 21CFR11.3(b)(6) as any combination of text, graphics, data, audio, pictorial, or other information representation in digital form that is created, modified, maintained, archived, retrieved, or distributed by an electronic system. This term applies specifically to records in electronic form that are created, modified, maintained, archived, retrieved, or transmitted, under any records requirements set forth in agency regulations. This also applies to electronic records submitted to the agency under requirements of the Federal Food, Drug, and Cosmetic Act and the Public Health Service Act, even if such records are not specifically identified in agency regulations.

### 5.3.8.3.1.2 SOURCE

An electronic record which is maintained by a regulated entity in an electronic system.

### 5.3.8.3.1.3 ORIGINAL COPY

A copy of a source electronic record provided during a regulatory activity. Original copies collected to support observations of potential violations or used as evidence in administrative or judiciary proceedings, including any original copy included in an EIR, memorandum, or C/R, must be stored as to maintain the chain of custody and assure the records may be verified any time after collection.

### 5.3.8.3.1.4 WORKING COPY

A copy of an electronic record which is created from the original copy and is used to review and analyze the records, decreasing the chance of unintentionally altering the original copy. This is an exact copy of the original copy electronic records.

### 5.3.8.3.2 - ELECTRONIC DATABASES AND QUERIES

Firms may use proprietary programs developed in-house or off the shelf programs to generate and/or store records used to show regulatory compliance, such as blood bank databases, drug production records, medical device complaints, and/or service records. These programs can often times be queried to generate electronic databases or summary data in a commonly used file format, such as Microsoft Excel. During an establishment inspection you may request and receive electronic databases or summary data generated by the firm from their databases. The methods used must maintain the integrity of the electronic data and prevent unauthorized changes. Do not
personally access a firm’s electronic records, databases, or source data during the course of an inspection.

When it is necessary to access a firm's data during an inspection:

1. Oversee the firm's personnel accessing their system and have them answer your questions.
2. Request the firm run queries specific to the information of interest.
3. Request the firm provide the parameters used to generate the data.
4. Request the firm to copy the data to electronic storage media.

Firm electronic data can be dynamic with real time updating. Your request may require the firm to develop one or more custom queries to provide the requested information. A custom report query is the method of using the reporting software to pull the specific data requested during the inspection (i.e., all complaints from the last 12 months with specific data fields). You must assume the query logic is not validated and take appropriate action to ensure the data is accurate and no data has been accidentally omitted due to a programming logic error occurring at the firm.

Reviewing data contained in electronic databases is generally most effectively accomplished with the use of a computer. Reviewing electronic data may require the transfer of electronic data to electronic storage media for you to use in your computer; see section 5.3.8.3.3 below for information on how to handle electronic storage media. Do not use the firm's equipment or personnel to perform repetitive queries or manipulation of the audited firm's own computerized data.

5.3.8.3.2.1 - REQUESTING COMPUTERIZED DATABASES

Before requesting a copy of computerized data, you should determine several things including information about the size and contents of the database, the program used by the firm, and the program you will use, among others. The following steps are useful in preparing for an electronic database request.

1. Determine the firm's application program used to maintain the data of interest. It is best to obtain data files in a format compatible with application programs currently used by the agency. Check the program you plan to use to ensure it can handle the file size you will be using.
2. You should determine what fields of information are routinely captured by the firm. This can be accomplished by requesting a printout of the data structure of the data file or observing the inputting of data at a computer terminal or workstation. It is common for databases to contain numbers or other coded information requiring translations from look up tables to give meaningful text. You should determine if information fields contain coded data, and if so, a code breakdown should be obtained. Information about code breakdowns should be located in the SOPs for that computerized system. Also be aware in relational databases, there may be linking data fields that exist in other tables that should also be considered in the overall data request.

3. If the files are too large to fit on electronic storage media, file compression can be used. If possible, ask that the firm prepare the data in a compression format that is self-extracting. Self-extracting files are executable files and should be virus scanned before and after executing. All electronic storage media should be scanned prior to being used on any FDA computer. Whatever compression utility is used, make sure you have the software to manipulate the files as needed.

5.3.8.3.3 - ELECTRONIC RECORDS RECEIVED ON ELECTRONIC STORAGE MEDIA

If you provide the electronic storage media to the firm, use only clean and preformatted media. An additional safeguard is to request the firm reformat the media on their own computer to assure it is usable and "clean". A universal serial bus (USB) flash drive cannot be used during foreign inspections.

Any request for electronic records on electronic storage media should be made with a computer application in mind and the information obtained should be useful. The electronic records should be in a format compatible with software applications knowledgeable to you and available from the Agency. Certain types of file conversion are difficult and should not be attempted without the necessary knowledge and availability of conversion type programs where applicable. Other file conversions are simple and have standard, built in conversion programs, such as converting a Microsoft Word document to PDF. If help is needed for file conversion, assistance may be available within the district, region or from the Office of Operations (OO).

Any electronic storage media containing electronic records received during the course of an inspection should be considered and handled as the original copy. The original copy of electronic records should be secured to assure the integrity of the data when used to support observations of potential violations or used as evidence in administrative or judiciary proceedings. Identify the original copy as an exhibit, place in a suitable container, e.g., FDA-525, and officially seal. Mark the FDA-525 or other container containing electronic storage media or other media and document the software type and version(s) required to
open the included software, e.g. Microsoft Word 2000, Microsoft Excel 2010, Windows Photo Viewer, etc. The electronic storage media or other media should be stored as part of the exhibits with the original EIR. See IOM 5.10.5.1.

There are no guarantees the files provided on electronic storage media will be usable data. It is your responsibility to make a working copy from each electronic storage media prior to closing the inspection. You will need to view the copied files and verify the files contain the information requested and the information is usable to you.

If you perform analysis, including sorts, pivot tables, or other reviews, on the working copy of an electronic database to develop or support observations, you should request the firm conduct the same analysis and provide a copy of this analysis. This can be done by requesting an electronic database that includes only the information of interest (e.g. an Excel spreadsheet of failures of a certain type for a specific time period), or requesting a paper copy of the information of interest. This will preclude or limit any errors that may have occurred from the investigator querying the database.

5.3.8.3.1 IDENTIFICATION AND SECURING ELECTRONIC STORAGE MEDIA

You should follow these steps to ensure proper identification and security of electronic storage media:

1. Label each original copy of electronic storage media
   - a. Firm name
   - b. Date and your initials
   - c. Initials by a representative of the firm (optional). If you provide the disk(s) to be used, use only new and preformatted disk(s).
   - d. The name of the appropriate software and version to ensure readability of the information.

2. Make a working copy of the electronic storage media
   - a. Virus scan the original storage media
   - b. Check the security on the storage media by viewing the “Security” tab under the “Properties” window (right click).
   - c. Copy the original of the electronic storage media.
   - d. Verify the data is usable.

3. Identify and place the original copy of electronic storage media in a Form FDA 525 or suitable container, label, and officially seal.

4. Prepare electronic record(s) for inclusion in the EIR, Memorandum, or C/R.
   - a. Include the applicable original and working copies with the EIR, Memorandum, or C/R.

5.3.8.5 - Listing of Records

If management requests a list of the copies of records you obtain, prepare it in duplicate and leave the original with the firm. Many firms prepare duplicate copies of documents requested during our inspections. In the interests of conserving inspectional time, you may ask the firm to prepare the list of copies concurrently with the photocopying and you then verify the accuracy. Do not use form FDA-484, Receipt for Samples. Describe the circumstances in your report including the name and title of the individual to whom you gave the list. Submit the duplicate list with your report as an exhibit.

5.3.8.6 - Patient and/or Consumer Identification on Records

During the course of many types of inspections and investigations you will review and collect records which specifically identify (by name) patients or consumers. Under most state Privacy Laws this information is confidential. Some firms we inspect may mistakenly believe this information is not releasable to the federal government. However, Federal laws preempt State laws; with few exceptions we are entitled to review and copy the complete record, including the identifying patient/consumer names. The Agency is then required to maintain the confidentiality of the records/files, as with any confidential record you collect. See IOM 5.10.5. Any disclosure of the information contained in the record(s) can only be by Law, i.e., judge's order, disclosure, Congressional order, etc. If you encounter resistance from the firm in providing patient records, you may refer them to 45 CFR 164.512(b) which explains the exemptions allowing FDA access to the patient records.

General, routine guidance is as follows:

1. For records copied as a result of injury or complaint investigation, where you obtain patient identification, the identification should remain intact and stored in the official FDA files. Frequently, medical releases must be obtained from a complainant, consumer or "next-of-kin". At least one or two extra should be obtained and stored in the files.

2. For any inspection/investigation involving a regulation required Informed Consent, such as clinical investigations, IRBs, bioequivalence testing, etc., patient identification should remain intact and stored in the official FDA files.

3. For most others, such as MQSA, plasmapheresis, blood donations, etc., only the patient initials and unique identifier supplied by the firm (such as donor number, donation number, etc.) need be routinely retained in the FDA files.

It is not uncommon for a firm to voluntarily purge the documents of the pertinent identifiers as they are copied. You must verify (by direct comparison to the original document) you received an accurate reproduction of the original, minus the agreed to purging, prior to accepting the copy.
As with any inspection there are times when the specific identifiers must be obtained, copied and retained, such as if/when further interview of the patient/consumer could be necessary. If in doubt, obtain the data. It is always easier to delete later than to return to obtain the information, especially in the few cases where questionable practices may result in the loss of the information.

All documents obtained containing confidential identifiers will be maintained as all documents obtained by FDA containing confidential information, i.e., in the official FDA files. Confidential identifiers may be flagged in the official FDA files for reference by reviewers to assure no confidential data are released under FOIA.

5.3.9 - REQUEST FOR SAMPLE COLLECTION

There are times one district will request another district to collect surveillance or compliance samples for it. The requesting district should provide as much of the following information as is available on specific shipments, using the FACTS Create Assignment Screen. See IOM Exhibit 5-9.

The following fields must be completed in order to save the assignment: Requesting Organization, Priority, Subject, POC Name, Op Code, Accomp Org, Num of Ops, and PAC. When you create a sample collection assignment, which will require laboratory analysis, you should also create an assignment for the laboratory, using operation 41.

The screen is organized in sections.

5.3.9.1 - FACTS Assignment Section

The Assignment section has the following fields:

Compliance Number: Enter the Compliance Number if known. This will make it easier to tie all associated activities together if the District is considering a compliance action. You can generate a compliance number after completing the mandatory fields on the Maintain Inspection Results screen.

Background Information: This is a free form field, which should briefly describe pertinent information related to the assignment.

ORA Reqd: This field only applies to assignments generated by Centers or other organizations outside of ORA. It will indicate whether or not ORA concurrence is required for the assignment.

ORA Cncrnc Num: This field is for the requesting organization (other than an ORA component) to indicate ORA concurrence for the assignment.

POC Name: This field indicates the point of contact in the requesting organization for the assignment.

Priority: Choose High or Routine

Remarks: This is a free form field, which should briefly describe the assignment.

Reporting Method: Indicate how the other district should notify the contact of problems with or status of the assignment. For example: e-mail, phone, etc.

Requesting Organization: Enter your District Office, if you are requesting a sample from another district or other appropriate FACTS organization.

Requestor Completion Date: Enter the completion date desired, using the format, MM/DD/YYYY.

Subject: Enter a subject for the assignment. It may be helpful to create a subject others will recognize as related to a specific action, for example a firm or product name.

5.3.9.2 - FACTS Operations Section

The Operations section has the following fields:

Estmtd Hours: Enter the number of hours you believe the assignment should take. This is done to assist the collecting district in planning their work.

Estmtd Smpl Cost: Enter the estimated sample cost, if known.

Op Code: Enter the operation code for the assignment. If you are requesting a sample collection, it is 31.

Requester Remarks: Enter as many details about the sample collection as you can. Include: date of shipment, number and size of units or amount, codes, carrier (routing and freight bill number), invoice number, and name of responsible firm with date of inspection (if one occurred).

Rqstr Prty: Enter High or Routine. This will default to the same data entered in the Assignment section if it was prepared first.

Subject: This will default to the same data entered in the Subject field in the Assignment section if it was prepared first.

Target Date: Enter the date to be completed by.

5.3.9.3 - FACTS Organizations Section

The Organizations section contains the following fields.

Accomp Org: Enter the District or other FACTS organization you are requesting collect the assignment. If you are completing the sample analysis assignment, be sure to enter a laboratory.

Num of Ops: Enter the number of sample collections or analyses you are requesting from the organization identified in the previous field.
Perf Org (Adhoc Work): If the performing organization is part of the accomplishing organization you are in, you may enter the performing organization here. If you are requesting the sample of another District, you will probably leave this blank.

The PACS & Products section of the form contains fields for entering the assignment PAC and Product code.

Enter the FEI number(s)/CFN(s) of the firm or firms from which the sample is to be collected in the Firms and Cross References section. See IOM 4.4.10.3.24.

5.3.10 - POST-INSPECTION NOTIFICATION LETTERS

Issuance of Post-inspection notification letters have been discontinued in all program areas. See FMD 145.

SUBCHAPTER 5.4 - FOOD

5.4.1 - FOOD INSPECTIONS

Food plant inspections are conducted to evaluate the methods, facilities, and controls used in manufacturing, storage and distribution of foods.

See CFSAN Office of Compliance's intranet website for the most current guidance (e.g., compliance programs, field assignments, field guidance).

5.4.1.1 - Preparation and References

Before undertaking an inspection:

1. Review the district files of the firm to be inspected and acquaint yourself with the firm's history, related firms, trademarks, practices and products. The review will identify products difficult to manufacture, require special handling, special processes or techniques, and hours of operation, which is especially important in bacteriological inspections. Remove, for subsequent investigations and discussion with management, Complaint/Injury Reports, which are marked for follow-up during the next inspection. See IOM 5.2.8.

2. Become familiar with current programs relating to the particular food or industry involved and relevant DFFPOI inspection guides. Become familiar with any applicable Compliance Policy Guide (CPG Chap 5).

3. Understand the nature of the assignment and whether it entails certain problems, e.g., Salmonella or other bacteriological aspects.

4. Review the FD&C Act Chapter IV - Food.

5. Review and become familiar with the appropriate parts of 21 CFR pertaining to foods, for example:
   a. 21 CFR Part 110 - GMP's on foods
   b. 21 CFR Parts 108 and 113 - Thermally Processed Low-Acid Foods Packaged in Hermetically Sealed Containers
   c. 21 CFR Part 114 - Acidified Foods
   d. 21 CFR Part 120 - HACCP Systems (covers Juice Processors)
   e. 21 CFR Part 123 - Fish and Fishery Products
   f. 21 CFR Part 129 - Processing and Bottling of Bottled Drinking Water
   g. 21 CFR Part 130, et al - Food Standards
   h. 21 CFR Part 1240 - Control of Communicable Disease
   i. 21 CFR Part 1250 - Interstate Conveyance Sanitation

6. Review reference materials on food technology and other subjects available in the District Inspectonal Reference Library.

7. If you are assigned to inspect food-service establishments under the FDA - Secret Service Agreement, you should use the most current copy of the "Food Code" and be standardized in its use. All Regional Food Service Specialists and most Interstate Travel Sanitation Specialists are standardized in use of the code.

8. Be familiar with the "Food Chemicals Codex". See IOM 5.4.4.3.

5.4.1.2 - Inspectonal Authority

See IOM subchapter 2.2 for broader information on this topic.

Authority to Obtain Records and Information in LACF and Acidified Foods Plants:

FDA's regulation in 21 CFR 113 requires commercial processors of low-acid foods packaged in hermetically sealed containers to maintain complete records of processing, production and initial distribution. 21 CFR 114 requires the same of commercial processors of acidified foods. 21 CFR 108.25(g) and 21 CFR 108.35(h) provide that a commercial processor shall permit the inspection and copying of the records required by 21 CFR 113 and 21 CFR 114 by duly authorized employees of FDA. The demand for these records must be in writing on an FDA 482a, Demand for Records, signed by you and must identify the records demanded.

5.4.1.2.1 - WRITTEN DEMAND FOR RECORDS

To obtain the records:

1. Prepare a FDA 482a, "Demand for Records", listing the records demanded. Describe the processing records to be reviewed and/or copied as accurately as you can, e.g., "All thermal process and production records mandated by 21 CFR 113 (or 114 if applicable) for the foods (state name of food) processed at this plant on (specific date or period of time)". If only a specific record is desired list it specifically as follows: e.g., "Fill Weight Records for #2 Filling Machine for the period 4-15-87 through 6-7-87."

2. Sign the form.

3. Issue the original to the same person to whom the FDA 482, "Notice of Inspection", was issued.

4. Submit an exact copy with your EIR.

5.4.1.2.2 - WRITTEN REQUEST FOR INFORMATION
4. Submit the carbon copy or exact copy with your EIR.
3. Issue the original to the same person to whom the FDA
2. Sign the form.
1. Prepare a FDA 482(b) - Request for Information listing
information:
the district to request it. If requested to obtain the
either request it directly from the processor or will direct
possible. If the process establishment data and
packaged in hermetically sealed containers shall provide
FDA with any information concerning processes and
procedures necessary by FDA to determine the adequacy
of the process. 21 CFR 108.25(c)(3)(ii) requires the same
of commercial processors of acidified foods. The
information in this regulation is the data on which the
processes are based. Many processors will not have this
information and in fact 21 CFR 113.83 requires only that
the person or organization establishing the process
permanently retain all records covering all aspects of
establishing the process. The processor should, however,
have in his files a letter or other written documentation
from a processing authority delineating the recommended
scheduled process and associated critical factors.

You may encounter situations where you believe control of
certain factors is critical to the process and there is no
evidence to document these factors were considered
when the process was established (e.g., a change in
formulation which could affect consistency). It is
appropriate to issue a written request for a letter or other
written documentation from a processing authority, which
delineates the recommended scheduled process and
associated critical factors. This represents the processing
authority's conclusions and should correlate with the filed
process.

If you believe control of certain factors is critical to the
process and are not delineated in the process authority's
recommendation or the filed process, obtain all available
information about the situation. Include the name of the
person or organization that established the process and
the specific practices of the firm. This information should
be included in your report and forwarded by your District
to the Center for Food Safety and Applied Nutrition,
Division of Enforcement (HFS-605) for review, as soon as
possible. If the process establishment data and
information is deemed necessary by the center, they will
either request it directly from the processor or will direct
the district to request it. If requested to obtain the
information:
1. Prepare a FDA 482(b) - Request for Information listing
the specific information requested. Specify each
product involved by food product name and form,
container size and processing method.
2. Sign the form.
3. Issue the original to the same person to whom the FDA
482, "Notice of Inspection", was issued.
4. Submit the carbon copy or exact copy with your EIR.

5.4.1.3 - Records Access Under Sections 414 and 704 of the FD&C Act

The Food Safety Modernization Act amended Section 414
of the Act to provide FDA with access and the ability to
copy records under the following circumstances:

1. FDA has a reasonable belief that an article of food, and
any other article of food that FDA reasonably believes
is likely to be affected in a similar manner:
   a. Is adulterated and presents a threat of serious
      adverse health consequences or death to humans
      or animals, and
   b. The records are needed to assist FDA in
determining whether the food is adulterated and
presents a threat of serious adverse health
consequences or death to humans or animals.

2. FDA believes that there is a reasonable probability that
use of or exposure to an article of food, and any other
article of food that the FDA reasonably believes is
likely to be affected in a similar manner:
   a. Will cause serious adverse health consequences or
defeat to humans or animals, and
   b. The records are needed to assist FDA in
determining whether there is a reasonable
probability that the use of exposure to the food will
cause serious adverse health consequences or
defeat to humans or animals.

If, during an inspection, you believe the above conditions
exist, and:
1. The firm refuses to provide access to the records, or
2. Based on past experience, the District anticipates that
   the firm may refuse to provide access to records, or
3. The firm requests FDA to provide a separate written
   request for records,

Notify your supervisor and consult with your district
Compliance Branch.

District management will obtain OE concurrence before
you issue the Form FDA 482c Notice of Inspection -
Request for Records. See Exhibit 5-10. District
management will notify FDA's Office of Emergency
Operations (OEO) of any situation requiring issuance of
Form FDA 482c. (OEO contact number: 1-866-300-4374
or 301-796-8240 - 24 hours/day.) OEO will notify CFSAN
or CVM, as appropriate, OO/OFFO, OE and OCC
according to standard operating procedures to obtain a
determination that the situation warrants issuance of Form
FDA 482c. OE, in consultation with CFSAN or CVM OEO,
OO/OFFO and the District, will determine if the standards
for records inspection in paragraphs (1) or (2) of section
414(a) have been met and identify the scope of the
records to request. Issue an FDA 482c, Notice of
Inspection – Request for Records. See Exhibit 5-10
according to their instructions.

FDA may at a later time, request additional records related
to the same article of food, or another article of food that is
likely to be affected in a similar manner, as long as the
criteria in 414(a)(1) or (a)(2) continue to be met. The
request for additional records may be verbal or written as
necessary to facilitate access to the records.

Investigators should document in the EIR a firm's refusal
to allow access to records or a firm's request for a written
request for records and issuance of Form FDA 482c.
5.4.1.4 - Food and Cosmetic Defense Inspectional Activities

Food and cosmetics security inspectional activities should be conducted during all routine food and cosmetics safety inspections. During the normal course of the inspection be alert to opportunities for improvement or enhancement of the firm’s food and cosmetics security preventive measures, as compared to those recommended in the guidance documents described below. You should not perform a comprehensive food and cosmetics security audit of the firm or conduct an extensive interview of management or employees in an attempt to determine the level of adoption of preventive measures listed in the guidance. The goal is to facilitate an exchange of information to heighten awareness on the subject of food and cosmetics security.

5.4.1.4.1 - FOOD AND COSMETIC SECURITY

Inspectional activities relative to food and cosmetic security for routine food and cosmetic establishment inspections should include:

1. Discussion with firm management of relevant FDA guidance documents including:
   a. Food Producers, Processors, and Transporters: Food Security Preventive Measures Guidance
   b. Importers and Filers: Food Security Preventive Measures Guidance
   c. Cosmetics Processors and Transporters: Cosmetics Security Preventive Measures Guidance
   d. Retail Food Stores and Food Service Establishments: Food Security Preventive Measures Guidance
   e. Dairy Farms, Bulk Milk Transporters, Bulk Milk Transfer Stations, and Fluid Milk Processors: Food Security Preventive Measures Guidance

   These documents should be used as references during inspections, as appropriate. If firm management does not already have a copy of the relevant guidance documents provide them with hard copies or information on how to obtain the guidance from FDA’s web site.

2. Identification of opportunities for improvement or enhancement of the firm’s food and cosmetic security preventive measures, as compared to those recommended in the guidance documents, and encouragement of management to make such improvements or enhancements to their security system.

   Keep in mind that: the guidance does not represent mandatory conditions or practices; some of the recommended food and cosmetics security preventive measures may not be appropriate or practical to the specific operation; and other means of achieving the goals of the preventive measures listed in the guidance may be more suitable for the specific operation than those cited as examples. The important message for management is to consider the goals of the food and cosmetics security preventive measures; evaluate the goals relative to the specifics of their operation; and address those that are relevant to the extent practical.

Food and cosmetics security observations should not be listed on form FDA-483, Inspectional Observations, unless they likewise constitute deviations from Current Good Manufacturing Practice. Security discussions should be handled discretely and should only involve management of the firm.

The fact that the discussion took place and, if applicable, that a copy of the guidance document(s) was provided should be recorded in the Summary section of the EIR. For example, under a section heading titled “Food and Cosmetics Security” you should only state, “A copy of the Food and Cosmetics Security Guidance documents were provided to and food and cosmetics security issues were discussed with (name of firm official).” The details of inspectional findings regarding security should NOT be recorded. You should also minimize the quantity and detail of notes taken relative to the firm’s food and cosmetics security program, taking only those needed to serve as a “memory jog” during the discussion with management.

5.4.1.4.2 - RECONCILIATION EXAMINATIONS

During routine food and cosmetic inspections, conduct one reconciliation examination during each food and cosmetic establishment inspection. The examinations are to be conducted on raw materials used in the manufacture of foods or cosmetics, or finished products received by the firm for further distribution. Preference should be given to products of foreign origin. Where possible, these examinations should be performed on products as they are received by the firm.

Consult the factory jacket for any information on special conditions in the facility that may affect selection of personal protective equipment; consult your supervisor for any recommendations on personal protective equipment; and have available all necessary personal protective equipment to conduct the activity.

As Part of an Import Field Examination and Entry Review - See IOM 6.3.1 and 6.4.3. For imported food and cosmetics, a reconciliation examination should be conducted:

1. Per Part A [IOM 5.4.1.4.3] during all routine import field exams. You should only report time under the Counter Terrorism PAC at the direction of your supervisor or if there is a for cause assignment.

2. In instances where review of entry information raises suspicion (resulting in a detailed reconciliation exam per Part B [IOM 5.4.1.4.4]).

A detailed reconciliation exam should be conducted when there are anomalies in entry declaration information. These may include new, unusual, or unfamiliar commodities, manufacturers, importers; suspicious trans- shipments; or credibility issues such as those between the product and declared country of origin.
If anomalies are found, entry documents should be requested and reviewed for discrepancies between the information declared through electronic filer submissions and that found in entry documents. Entry documents may include invoices, bills of lading, export certifications, and other relevant documents obtained from the importer, filer, or manufacturer/processor of the product. Fields in which discrepancies are found that may raise concern include country of origin, manufacturer, product description, product code, and quantity.

Avoid duplication of examination of the same foreign manufacturer, unless a prior reconciliation examination disclosed an unexplained discrepancy.

Follow guidance in IOM 5.4.1.4.3 to IOM 5.4.1.4.4 below for domestic and import reconciliation exams.

5.4.1.4.3 - RECONCILIATION EXAMINATION GUIDANCE PART A

Reconciliation examinations are performed to ensure that:
1. The food or cosmetic is what it purports to be
2. There are not unexplained differences in the quantity of product ordered, shipped, and received, and
3. There are no signs of tampering or counterfeiting.

Before initiating the exam make a general assessment of the appearance of the lot. Look for packaging that: appears to have been opened and resealed; appears wet, stained, punctured, or powdered. Also be alert to abnormal chemical odors. If any of these conditions are detected stop the exam and contact your supervisor for guidance. If the lot appears normal proceed with the examination. To the extent possible the exam should be performed in a well-ventilated, well-lit area.

Determine, to the extent possible, whether:
1. The actual goods in a lot are the same as those that are declared in the shipping documents
2. There is consistency in the manufacturer declared on the product labeling, bulk product packaging, and shipping documents; and
3. There is no (unexplainable) inconsistency in actual quantity of goods in the lot, and the quantity ordered and declared in the shipping documents.

If no unexplained inconsistencies are detected, no further action is indicated.

If unexplainable inconsistencies are detected, document the occurrence, including photographs of the labeling and packaging, and an accurate count of the lot. Contact your supervisor, who should, in the case of imported products, contact the U.S. Customs and Border Protection for appropriate action. If the examination discloses evidence that inaccurate product identification data was submitted to the OASIS entry screening system, the District should evaluate the need for follow-up with a compliance filer evaluation and consider providing the information to the U.S. Customs and Border Protection for appropriate action.

In addition, if unexplained inconsistencies are detected, follow part B [IOM 5.4.1.4.4] of this guidance while conducting a detailed reconciliation exam.

5.4.1.4.4 - RECONCILIATION EXAMINATION GUIDANCE PART B

Open the shipping packaging of a quantity of product approximating the square root of the number of shipping cartons/packages in the lot, and examine the contents. Look for the following:
1. Product identity on the package that does not match the identity declared on the shipping documents
2. Mixed product sizes within a carton or within the lot;
3. Product sizes that do not match the sizes declared on the shipping documents
4. Differences in product configuration or package type (e.g. plastic containers mixed with glass jars or aluminum or steel cans)
5. Easily apparent variations in weight
6. Product labels that display crude, unprofessional, or inconsistent styles of print, color or use of language
7. Unusual placement of labels (e.g. off-center)
8. Variations in lot coding ink color, appearance of embossing, or format (e.g., two line vs. three line, use of letters numbers and symbols), unusually excessive use of a single code in a very large lot
9. Differences between the actual can codes in the lot and those listed on the shipping documents
10. The existence of a tamper-evident notice on the labeling when the packaging does not contain a tamper-evident feature
11. Product that is beyond its expiration date
12. Inconsistencies in expiration dates within a lot

If no unexplainable discrepancies are noted select at least 1 package at random from the entire shipment and examine their contents. For those products that the contents are visible through the package it is not necessary to open the package. For other products, open the package and examine and field destroy the contents. Look for the following:
1. Differences between the product and that which is declared on the label
2. Color differences in the product between containers of the same lot
3. Style differences in the product between containers of the same lot or between the actual product and the label and document declaration (e.g., sliced vs. whole, colorless noodles vs. egg noodles)
4. Readily detectable abnormal odors (e.g. strong decomposition, bitter almond, petroleum odor, garlic, chlorine, sulfur). Note: specific sensory examination is not expected.

Verification that the product is consistent with the product ordered may require that you obtain information from the owner of the goods, importer, filer, or custom house broker. Review of the following types of documentation may be necessary to accomplish the above instructions, to the extent that they are available: authentic label supplied by the owner of the goods, importer, filer, or custom house broker.
registrant is issued a system generated 11 digit registration number. Upon completion, the registration information is maintained in the Unified Registration and Listing System (FURLS) database. A facility is not registered until all required fields have been completed in FFRM. 

When performing an establishment inspection or reconciliation examination, follow these instructions:

1. If there are no signs of tampering or counterfeiting, use level I protection, which consists of: work gloves; coveralls; work boots; and in a dusty situation, a dust mask.
2. If there are signs of tampering or counterfeiting, use level II protection and consult your supervisor for any additional safety precautions needed. Level II protection consists of: work gloves worn over surgical gloves; full face respirator with appropriate cartridges; disposable coveralls; and work boots.

5.4.1.5 - Food Registration

Section 415 of the FD&C Act (21 U.S.C. 350d) requires most domestic and foreign facilities that manufacture/process, pack, or hold food for human or animal consumption in the United States to register with FDA before operations commence. Section 415 also requires food facilities to renew their registration biennially. FDA requires renewals to be submitted between October 1 and December 31 of each even-numbered year. Facilities may register electronically at http://www.access.fda.gov, by mail, or by CD-ROM for multiple submissions, to Food and Drug Administration, Food Facility Registration, HFS-651, 5100 Paint Branch Parkway, College Park, MD, 20993-2804. FDA maintains the registration information in the Food Facility Registration Module (FFRM) within the FDA Unified Registration and Listing System (FURLS) database. A facility is not registered until all required fields have been completed in FFRM. Upon completion, the registrant is issued a system generated 11 digit registration number.

For food facilities that are required to register, the owner, operator, or agent in charge of a facility must provide the following:

1. Facility name, address, phone number, and emergency contact phone number;
2. Parent company name, address, and phone number (if applicable);
3. Name, address, and phone number of the owner, operator, or agent in charge;
4. Email address for the contact person of the facility or, in the case of a foreign facility, the U.S. agent for the facility;
5. All trade names the facility uses;
6. Applicable food product categories, as listed on the registration form;
7. Name, address, and phone number of a foreign facility’s U.S. agent and phone number of the facility’s emergency contact if it is someone other than the U.S. agent;
8. Certification that the information submitted is true and accurate and that the person submitting the registration is authorized to do so; and
9. Assurance that FDA will be permitted to inspect the facility at the times and in the manner permitted by the FD&C Act (section 415(a)(2)).

Section 415(b) of the FD&C Act also provides FDA with authority to suspend the registration of a facility when:

1. FDA determines that food manufactured, processed, packed, received, or held by a registered facility has a reasonable probability of causing serious adverse health consequences or death to humans or animals (SAHCODHA); and
2. That facility:
   a. Created, caused, or was otherwise responsible for that reasonable probability of SAHCODHA; or
   b. Knew of, or had reason to know of, the reasonable probability of SAHCODHA, and packed, received, or held such food.

The purpose of registration is to provide FDA with sufficient and reliable information about food facilities. Registration will help provide information on the origin and distribution of food that may be associated with a real and potential threat to public health. In the event of a foodborne outbreak of illness, registration information will enable FDA to identify the food facility representatives and to investigate the source and cause of the outbreak. It will also enable FDA to identify and contact other facilities that might be associated with the food causing the outbreak.

Under section 301(dd) of the FD&C Act (21 U.S.C. 331(dd)), the failure to register a food facility is a prohibited act. Food from a foreign facility that is not registered may be held at the port of entry (section 301(l)) of the FD&C Act (21 U.S.C. 381(l)).

5.4.1.5.1 - Facilities Exempted from Registration

The following food facilities do not have to register (21 CFR 1.226):

1. A foreign facility, if food from such facility undergoes further manufacturing/processing
(including packaging) by another facility outside the U.S. A foreign facility is not exempt under this provision if the further manufacturing/processing (including packaging) conducted by the subsequent facility consists of adding labeling or any similar activity of a de minimis nature. The facility conducting the de minimis activity also must register.

2. Farms that are devoted to the growing and harvesting of crops, the raising of animals (including seafood), or both. Washing, trimming of outer leaves of, and cooling produce are considered part of harvesting. The term “farm” includes:
   a. Facilities that pack or hold food, provided that all food used in such activities is grown, raised, or consumed on that farm or another farm under the same ownership; and
   b. Facilities that manufacture/process food, provided that all food used in such activities is consumed on that farm or another farm under the same ownership.

3. Retail food establishments whose sales to consumers exceed their sales to non-consumers (businesses are considered non-consumers).

4. Restaurants that prepare and serve food directly to consumers for immediate consumption.

5. Nonprofit food establishments in which food is prepared for, or served directly to, the consumer.

6. Fishing vessels, including those that not only harvest and transport fish but also engage in practices such as heading, eviscerating, or freezing intended solely to prepare fish for holding on board a harvest vessel. However, those fishing vessels that otherwise engage in processing fish are required to register. For the purposes of this section, “processing” means handling, storing, preparing, shucking, changing into different market forms, manufacturing, preserving, packing, labeling, dockside unloading, holding, or heading, eviscerating, or freezing other than solely to prepare fish for holding on board a harvest vessel. However, those fishing vessels that otherwise engage in processing fish are required to register. For the purposes of this section, “processing” means handling, storing, preparing, shucking, changing into different market forms, manufacturing, preserving, packing, labeling, dockside unloading, holding, or heading, eviscerating, or freezing other than solely to prepare fish for holding on board a harvest vessel.


8. Other exemptions from registration in the final rule are based on the definition of food included within the scope of the registration regulation. Facilities that manufacture/process, pack, or hold food contact substances (including packaging materials) (21 CFR 1.227(b)(4)(i)(A)) or pesticides (21 CFR 1.227(b)(4)(i)(B)) are exempt from registration.

5.4.1.5.2 - AGENCY WEBSITE LINK

Additional information relating to food facility registration is available at the following website: http://www.fda.gov/Food/GuidanceRegulation/FoodFacilityRegistration/default.htm.

5.4.1.5.3 - INSPECTIONAL GUIDANCE

See the guidance in Compliance Policy Guide Sec. 110.300 Registration of Food Facilities Under the Public Health Security and Bioterrorism Preparedness and Response Act of 2002. During inspection of a domestic or foreign facility that is required to register, make sure that firm’s management is aware of the food facility registration requirements. Inform the firm’s management that information regarding food facility registration and penalties for failure to register is available at the following website: http://www.fda.gov/Food/GuidanceRegulation/FoodFacilityRegistration/default.htm. For facilities that are required to register, but have not done so, encourage electronic registration and provide them with the web site address for electronic registration http://www.access.fda.gov. If the firm needs to submit the hard copy registration form, inform them that they may obtain a registration form to complete and submit by mail at http://www.fda.gov/downloads/AboutFDA/ReportsManualsForms/Forms/UCM071977.pdf). Also encourage the firm to submit the optional information on the registration form to assist and facilitate FDA’s future communications with the firm.

Document the registration status of the firm and the registration discussion with the firm’s management in the “Summary of Findings and Discussion with Management” section of the EIR. If you obtain the firm’s food facility registration number, do not record the registration number in the EIR. Do not include observations about a firm’s failure to register on the Form FDA 483.

If the registration information obtained during the inspection is different from the information in FFRM, send an email to CFSANFoodFacilityRegistration@fda.hhs.gov with the facility name, FEI or registration number, and the specific registration information that is inaccurate (e.g., type of activity, facility name/address, emergency contact information). If the facility is operating with no registration, suspended registration, invalid registration, or a cancelled registration send an email to CFSANFoodFacilityRegistration@fda.hhs.gov describing the situation with the facility name and FEI and denote this in the EIR.

5.4.1.6 - CFSAN Bio-research Monitoring
Bio-research monitoring (BIMO) assignments for foods will generally be issued by the Center for Food Safety and Applied Nutrition (CFSAN) (see IOM 5.5.6).

5.4.2 - PERSONNEL

5.4.2.1 - Management

Follow the guidance described in IOM 5.3.6 when documenting individual responsibility including obtaining the full name and titles of the following individuals:
1. Owners, partners, or officers.
2. Other management officials or individuals supplying information.
3. Individuals to whom credentials were shown and FDA 482, Notice of Inspection, and other inspectional forms issued.
4. Individuals refusing to supply information or permit inspection.
5. Individuals with whom inspectional findings were discussed or recommendations made.

Regulations require plant management take all reasonable measures and precautions to assure control of communicable disease, employee cleanliness, appropriate training of key personnel, and compliance by all personnel with all requirements of 21 CFR 110.10, 113.10, and 114.10.

Determine if adequate supervision is provided for critical operations where violations are likely to occur if tasks are improperly performed.

5.4.2.2 - Employees

Improper employee habits may contribute to violative practices in an otherwise satisfactory plant. Observe the attitude and actions of employees during all phases of the inspection. Observe employees at their work stations and determine their duties or work functions. Note whether employees are neatly and cleanly dressed and whether they wear head coverings which properly cover their hair.

Determine if employees working with the product have obvious colds, or infected sores, cuts, etc. Under no circumstance should you swab a sore, touch or remove a bandage from an employee in an attempt to obtain bacteriological data. To do so is a violation of personal privacy, possibly hazardous to you and/or the employee, and usually provides little useful data.

Note whether employees eat while on duty.

Observe and record insanitary employee practices or actions showing employees handling or touching insanitary or dirty surfaces and then contacting food products or direct food contact surfaces. Such practices might include employees spitting, handling garbage, placing their hands in or near their mouths, cleaning drains, handling dirty containers, etc. and then handling food product without washing and sanitizing their hands.

Observe whether employees comply with plant rules such as, "No smoking", "Keep doors closed", "Wash hands before returning to work", etc. See IOM 5.4.7.2.2.

Be alert to employees handling insanitary objects, then quickly dipping their hands in sanitizing solutions without first washing them. Depending upon the amount and type of filth deposited on the hands during the handling of insanitary objects, such attempts at sanitizing are questionable at best. Sanitizers work most effectively on hands, which have been first cleaned by washing with soap and water.

Conversations with employees doing the work may provide information on both current and past objectionable practices, conditions and circumstances. These should be recorded in your notes.

Where appropriate, determine employee education and training. Also determine type, duration, and adequacy of firm's training programs, if any, to prepare employees for their positions and to maintain their skills.

5.4.3 - PLANTS AND GROUNDS

Observe the general nature of the neighborhood in which the firm is located. Environmental factors such as proximity to swamps, rivers, wharves, city dumps, etc., may contribute to rodent, bird, insect or other sanitation problems.

5.4.3.1 - Plant Construction, Design and Maintenance

Determine the approximate size and type of building housing the firm and if suitable in size, construction, and design to facilitate maintenance and sanitary operations. Check placement of equipment, storage of materials, lighting, ventilation, and placement of partitions and screening to eliminate product contamination by bacteria, birds, vermin, etc. Determine any construction defects or other conditions such as broken windows, cracked floor boards, sagging doors, etc. which may permit animal entry or harborage.

Inspect toilet facilities for cleanliness, adequate supplies of toilet paper, soap, towels, hot and cold water, and hand washing signs. Check if hand washing facilities are hidden, or if located where supervisory personnel can police hand washing.

Determine who is responsible for buildings and grounds maintenance. Many facilities such as docks, wharves, or other premises are owned and maintained by other firms, municipalities, or individuals for lease for manufacturing operations. Determine who is legally responsible for repairs, maintenance, rodent proofing, screening, etc. Evaluate the firm's attitude toward maintenance and cleaning operations.

5.4.3.2 - Waste Disposal
Waste and garbage disposal poses a problem in all food plants depending upon plant location and municipal facilities available.

Check the effectiveness of waste disposal on the premises and ensure it does not cause violative conditions or contribute toward contamination of the finished products. Check for in-plant contamination of equipment and/or product, if its water is supplied from nearby streams, springs, lakes or wells.

Suspected dumping of sewage effluent into nearby streams, lakes, or bay waters near water intakes can be documented by color photographs and water soluble fluorescein sodium dye. Place approximately two ounces dye, which yields a yellowish red color, into the firm's waste system and/or toilets, as applicable, and flush the system. The discharge area of the effluent becomes readily visible by a yellowish-red color on the surface of the water as the dye reaches it. Color photographs should be taken.

Determine collecting or flushing methods used to remove waste from operating areas. If water is used, determine if it is recirculated and thus may contaminate equipment or materials.

Determine the disposition of waste materials that should not be used as human food such as rancid nuts, juice from decomposed tomatoes, etc.

Determine the disposition of waste, garbage, etc., which contain pesticide residues. Determine how this is segregated from waste material which contains no residues and which may be used for animal feed.

5.4.3.3 - Plant Services

If applicable, check steam generators for capacity and demand. Demand may reach or exceed the rated capacity, which could affect adequacy of the process. Check boiler water additives if steam comes in direct contact with foods.

Check central compressed air supply for effective removal of moisture (condensate) and oil. Determine if any undrained loops in the supply line exist where condensate can accumulate and become contaminated with foreign material or microorganisms.

5.4.4 - RAW MATERIALS

List in a general way the nature of raw materials on hand. Itemize and describe those, which are unusual to you, or involved in a suspected violation (copy quantity of contents and ingredient statements, codes, name of manufacturer or distributor, etc.). Be alert for additives and preservatives. Evaluate the storage of materials. Determine the general storage pattern, stock rotation and general housekeeping. Materials should be stored so they are accessible for inspection. Thoroughly check ceilings, walls, ledges, and floors in raw material storage areas for evidence or rodent or insect infestation, water dripping or other adverse conditions.

5.4.4.1 - Handling Procedure

Determine if growing conditions relative to disease, insects, and weather are affecting the raw material. Check measures taken for protection against insect or rodent damage. Raw materials may be susceptible to decomposition, bruising or damage, e.g., soft vegetables and fruits delivered in truckload lots. Determine the holding times of materials subject to progressive decomposition.

5.4.4.2 - Condition

Evaluate the firm's acceptance examination and inspection practices including washing and disposition of rejected lots. Where indicated, examine rejected lots and collect appropriate samples and report consignees.

Determine the general acceptability of raw materials for their intended use and their effect on the finished product. Raw stocks of fruits or vegetables may contribute decomposed or filthy material to the finished product. Be alert for use of low quality or salvage raw materials. Check bags, bales, cases and other types of raw material containers to determine signs of abnormal conditions, indicating presence of filthy, putrid or decomposed items. Check any indication of gnawed or otherwise damaged containers, to ascertain if material is violative. Be alert to contamination of raw materials by infested or contaminated railroad cars or other carriers.

Document by photographs, exhibits or sketches any instances where insanitary storage or handling conditions exist.

5.4.4.3 - Food Chemicals Codex

Any substance used in foods must be food-grade quality. FDA regards the applicable specifications in the current edition of the publication "Food Chemicals Codex" as establishing food-grade unless FDA publishes other specifications in the Federal Register.

Determine whether firm is aware of this publication and whether or not they comply.

5.4.5 - EQUIPMENT AND UTENSILS

By arriving before processing begins, you are able to evaluate conditions and practices not otherwise observable before plant start-up. This includes adequacy of clean-up, where and how equipment is stored while not in use, how hand sanitizing solutions and food batches are prepared and if personnel sanitize their hands and equipment before beginning work.

Dirty or improperly cleaned equipment and utensils may be the focal point for filth or bacterial contamination of the finished product. Examine all equipment for suitability and
accessibility for cleaning. Determine if equipment is constructed or covered to protect contents from dust and environmental contamination. Open inspection ports to check inside only when this can be done safely. Notice whether inspection ports have been painted over or permanently sealed.

5.4.5.1 - Filtering Systems

Observe the firm's filtering systems and evaluate the cleaning methods (or replacement intervals of disposable filters) and schedules. Check types of filters used. There have been instances where firms have relied on household furnace type filters.

5.4.5.2 - Sanitation of Machinery

Check the sanitary condition of all machinery. Determine if equipment is cleaned prior to each use and the method of cleaning. If the firm rents or leases equipment on a short-term basis, report prior cleaning procedures. Equipment may have been used for pesticides, chemicals, drugs, etc., prior to being installed and could therefore be a source of cross-contamination.

5.4.5.3 - Conveyor Belt Conditions

Inspect conveyor belts for build-up of residual materials and pockets of residue in corners and under belts. Look in inspection ports and hard-to-reach places inside, around, underneath, and behind equipment and machinery for evidence of filth, insects, and/or rodent contamination. Chutes and conveyor ducts may appear satisfactory, but a rap on them with the heel of your hand or a rubber mallet may dislodge static material, which can be examined. See IOM 4.3.7.7.3 for procedure on taking In-line Sample Subs.

5.4.5.4 - Utensils

Determine how brushes, scrapers, brooms, and other items used during processing or on product contact surfaces are cleaned, sanitized and stored. Evaluate the effectiveness of the practices observed.

5.4.5.5 - Mercury and Glass Contamination

Be alert for improper placement or inadequately protected mercury switches, mercury thermometers, or electric bulbs. Breakage of these could spray mercury and glass particles onto materials or into processing machinery.

5.4.5.6 - UV Lamps

If firm is using ultra violet (UV) lamps for bacteria control, check if it has and uses any method or meters to check the strength of UV emissions. If so, obtain methods, procedures, type equipment used, and schedule for replacement of weak UV bulbs.

5.4.5.7 - Chlorine Solution Pipes

In plants where chlorine solution is piped, check on type of pipe used. Fiberglass reinforced epoxy pipe has been observed to erode inside through the action of the chlorine solution. This poses a threat of contamination from exposed glass fibers. Pipes made with polyester resin do not deteriorate from this solution.

5.4.5.8 - Sanitation Practices

Observe sanitizing practices throughout the plant and evaluate their effectiveness, degree of supervision exercised, strength, time, and methods of use of sanitizing agents. Determine the use, or absence of, sanitizing solutions both for sanitizing equipment and utensils as well as for hand dipping. If chlorine is used, 50 ppm - 200 ppm should be used for equipment and utensils, while a 100 ppm will suffice for hand dipping solutions. Sanitizing solutions rapidly lose strength with the addition of organic material. The strength of the solution should be checked several times during the inspection.

5.4.6 - MANUFACTURING PROCESS

Where helpful to describe equipment and processes, draw flow plans or diagrams to show movement of materials through the plant. Generally a brief description of each step in the process is sufficient. List all quality control activities for each step in the process and identify Critical Control Points. Provide a full description when necessary to describe and document objectionable conditions, or where the assignment specifically requests it.

Observe whether hands and equipment are washed or sanitized after contact with insanitary surfaces. For example:

1. Workers do general work, then handle the product;
2. Containers contact the floor, then are nested or otherwise contact product or table surfaces;
3. Workers use common or dirty cloths or clothing for wiping hands;
4. Product falls on a dirty floor or a floor subject to outside foot traffic and is returned to the production line.

Be alert for optimum moisture, time and temperature conditions conducive to bacterial growth.

In industries where scrap portions of the product are re-used or re-worked into the process (e.g., candy and macaroni products), observe the methods used in the re-working and evaluate from a bacteriological standpoint. Re-working procedures such as soaking of macaroni or noodle scrap to soften or hand kneading of scrap material offers an excellent seeding medium for bacteria.

When a product is processed in a manner which destroys micro-organisms, note whether there are any routes of
recontamination from the "raw" to the processed product (e.g. dusts, common equipment, hands, flies, etc.).

5.4.6.1 - Ingredient Handling

Observe the method of adding ingredients to the process. Filth may be added into the process stream from dust, rodent excreta pellets, debris, etc. adhering to the surface of ingredient containers. Evaluate the effectiveness of cleaning and inspectional operations performed on the materials prior to or while adding to the process. Determine specific trimming or sorting operations on low quality or questionable material. Observe and report any significant lags during the process or between completion of final process and final shipping. For example, excessive delay between packing and freezing may be a factor in production of a violative product.

5.4.6.2 - Formulas

The Act does not specifically require management to furnish formula information except for human drugs, restricted devices and infant formulas. Nonetheless, they should be requested especially when necessary to document violations of standards, labeling, or color and food additives. Management may provide the qualitative formula but refuse the quantitative formula.

If formula information is refused, attempt to reconstruct formula by observing:
1. Product in production,
2. Batch cards or formula sheets,
3. Raw materials and their location.

Any refusal to furnish requested information is reported in your EIR under the refusal heading.

5.4.6.3 - Food Additives

Refer to the food additives programs in CP (Chapter 9) for instructions on conducting establishment inspections of firms manufacturing food additive chemicals. Information is also available in ORA's "Guide to Inspections of Manufacturers of Miscellaneous Food Products - Volume II.

When making food plant inspections direct your evaluation of food additives only to those instances of significant violation or gross misuse.

Routine inspectional coverage will be directed primarily to the following two types of additives:
1. Unauthorized and illegal as listed in the Food Additive Status List (safrole, thiourea, et al), and
2. Restricted as to amount in finished food.

Because of special problems, exclude the following additives from coverage during routine inspections:
1. Packaging materials,
2. Waxes and chemicals applied to fresh fruit and vegetables,
3. Synthetic flavors and flavoring components except those banned by regulations or policy statements (these products will be covered under other programs), and
4. Food additives in feeds (these products will be covered under other programs).

The Food Additives Status List (FASL) found on the CFSAN website contains an alphabetical listing of substances, which may be added directly to foods or feeds and their status under the Food Additives Amendment and Food Standards. In addition, a few unauthorized or illegal substances are included.

You may encounter substances not included in the Food Additives Status List (FASL). Such substances will include:
1. Safe substances not on the list of items Generally Recognized as Safe (GRAS) which are not published in the regulations, i.e., salt, cane sugar, corn syrup, vinegar, etc.;
2. Synthetic flavoring substances because of their indefinite status;
3. Substances pending administrative determination,
4. Substances granted prior sanction for specific use prior to enactment of the Food Additives Amendment.

Give primary attention to unauthorized substances. Document and calculate levels of restricted-use additives in finished food only where gross misuse or program violations are suspected as follows:
1. List ingredients, which may be restricted substances or food additives, and determine their status by referring to the current FASL. Report complete labeling on containers of these substances.
2. Obtain the quantitative formula for the finished product in question.
3. Determine the total batch weight by converting all ingredients to common units.
4. Calculate the theoretical levels in the final product of all restricted or unauthorized ingredients from the formula by using the Food Additives Nomographs. See IOM Exhibit 5-11.
5. Determine probable level of restricted ingredients by observing the weight of each ingredient actually put into the batch.

5.4.6.4 - Color Additives

Evaluate the status of color additives observed during each establishment inspection by using the Color Additive Status List and the Summary of Color Additives Listed in the United States in Food, Drugs, Cosmetics, and Medical Devices. Both of these links can be found on the CFSAN website. These lists provide the current status and use limitations of most color additives likely to be found in food, drug, device, or cosmetic establishments.

Stocks of delisted and uncertified colors may be found in the possession of manufacturers where there is no evidence of misuse. Advise the firm of the status of these colors. If management wishes to voluntarily destroy such
colors, witness the destruction and include the facts in your EIR. If the firm declines to destroy the colors, determine what disposition is planned, i.e., use in non-food, non-drug, non-cosmetic or non-medical device products. The validity of certification information can be checked by accessing the online Color Certification Database system maintained by the Office of Cosmetics and Colors or contact Ray Decker, Director, Division of Color Certification and Technology, HFS-105, by e-mail at raymond.decker@fda.hhs.gov to be granted user privilege.

Where decertified or restricted-use colors are used in manufacturing food, drug, device, or cosmetics products, proceed as follows:

1. Collect an Official Sample consisting of the color and the article in which it is being used. Make every effort to collect interstate shipments of the adulterated product before attempting to develop a 301(k) or 301(a) case. When regulatory action is an alternative, obtain sufficient interstate records to cover both the color and the basic ingredients of the manufactured product. Refer to IOM Sample Schedule, Chart 9 - Sampling Schedule for Color Containing Products for guidance.

2. Document the use of decertified colors after the decertifying date. Documentation should include batch cards, employee statements, code marks indicating date of manufacture, color certification number, etc. The presence of color in the finished product will be confirmed by your servicing laboratory.

5.4.6.5 - Quality Control

The objective of quality control is to ensure the maintenance of proper standards in manufactured goods, especially by periodic random inspection of the product. Your inspection should determine if the firm's quality control system accomplishes its intended purpose. Establish responsibility for specific operations in the control system. Determine which controls are critical for the safety of the finished product.

5.4.6.5.1 - INSPECTION SYSTEM

Determine what inspectional control is exercised over both raw materials and the processing steps. Such inspection may vary from simple visual or other organoleptic examination to elaborate mechanical manipulation. Determine what inspection equipment is used, i.e., inspection belts, sorting belts, grading tables, ultraviolet lights, etc. Ascertain its effectiveness, maintenance or adjustment schedules. Where indicated, determine the name of the manufacturer of any mechanical inspection device and the principles of its operation.

Evaluate the effectiveness of the personnel assigned to inspection operations. Determine if the inspection belts or pick-out stations are adequately staffed and supervised.

Determine the disposition of waste materials, which are unfit for food or feed purposes.

5.4.6.5.2 - LABORATORY TESTS

Describe routine tests or examinations performed by the firm's laboratory and the records maintained by the firm. Determine what equipment is available in the laboratory and if it is adequate for the purpose intended. If the firm uses a consulting laboratory, determine what tests are performed and how often. Review laboratory records for the period immediately preceding the inspection.

5.4.6.5.3 - MANUFACTURING CODE SYSTEM

Obtain a complete description of the coding system with any necessary keys for interpretation. Provide an example by illustrating the code being used at the time of the inspection. (See 21 CFR 113.60(c) and 114.80(b)). Report coding systems, which require the use of ultra-violet light for visibility. Hermetically sealed containers of low acid processed food must be coded in a manner clearly visible. (See 21 CFR 113.60). Check 21 CFR 113 and 114 for regulations on coding for the type plant you are inspecting.

5.4.6.6 - Packaging and Labeling

Evaluate storage of packaging materials including protection from contamination by rodents, insects, toxic chemicals or other materials. Appraise the manner in which containers are handled and delivered to the filling areas. Determine if there is likelihood of chipping of glass or denting, puncturing, tearing, etc., of packaging materials. Observe the preparation of containers prior to filling. Consider any washing, steaming, or other cleaning process for effectiveness. Determine, in detail, the use of air pressure or other cleaning devices.

5.4.6.6.1 - QUANTITY OF CONTENTS

If slack fill is suspected, weigh a representative number of finished packages. See IOM 4.3.8 for net weight procedure. Sets of official weights are available in the district servicing laboratory. These may be used to check the accuracy of firm's weighing equipment.

5.4.6.6.2 - LABELING

Check the sanitary condition of labelers and equipment feeding cans to, and away from, the labeler. Determine if old product is present on any equipment which touches the can end seams, in the presence of moisture carry-over from the can cooling operation. Check availability of floor drains in the labeling area. Absence of floor drains could indicate infrequent cleaning of the equipment unless it is physically moved to another area for cleaning.

Determine what labels are used and what labeling is prepared or used to accompany or promote the product. Obtain specimens of representative labels and labeling including pamphlets, booklets, and other promotional material.
5.4.6.3 – NUTRITIONAL AND ALLERGEN LABELING


5.4.7 - SANITATION

Documented observation of the conditions under which food products are processed, packed, or stored is essential to the proper evaluation of the firm’s compliance with the law. This involves the determination of whether or not insanitary conditions contribute to the product being adulterated with filth, rendered injurious to health, or whether it consists in whole or in part of a filthy, putrid or decomposed substance.

Observations that dirt, decomposed materials, feces or other filthy materials are present in the facility and there is a reasonable possibility these filthy materials will be incorporated in the food are also ways of determining products may have become contaminated.

5.4.7.1 - Routes of Contamination

It is not sufficient to document only the existence of insanitary or filthy conditions. You must also demonstrate how these conditions contribute or may contribute to contaminating the finished product. Investigate and trace potential routes of contamination and observe all means by which filth or hazardous substance may be incorporated into the finished product. For example, defiled molding starch in a candy plant may contribute filth to candy passing through it, or filth in insect or rodent contaminated raw materials may carry over into the finished product. IOM Section 4.3.7 contains instructions on sample collection techniques for adulteration violations, including instructions for field exams and sample collections to document evidence of rodent, insect, etc., contaminated lots, and instructions for in-line sampling, including bacteriological samples. Finished product sample sizes for filth and micro collections can be found in the applicable Compliance Program (CP) or DFI "Guide to Inspections of ***."

5.4.7.1.1 - INSECTS

Insect contamination of the finished product may result from insect infested raw material, infested processing equipment or insanitary practices, and by insanitary handling of the finished product. When routes of contamination with insect filth are encountered, identify the insects generally, e.g., weevils, beetles, moths, etc. If qualified, identify as to species. You must be correct in your identification. See IOM Appendix A.

5.4.7.1.2 - RODENTS

Rodent contamination of the finished product may result from using rodent defiled raw materials, exposure to rodents during processing, and by rodent depredation of the finished product. When evidence of rodents are discovered, you should thoroughly describe its composition, quantity, estimated age and location. Explain its significance and potential for product contamination. See IOM Section 4.3.7.4.2.3 – Summary of Sample for Rodent Evidence.

5.4.7.1.3 - PESTICIDES

Pesticide contamination of the finished product may be the result of mishandling of food products at any stage in manufacturing or storage. The use of toxic rodenticides or insecticides in a manner, which may result in contamination, constitutes an insanitary condition. Where careless use of these toxic chemicals is observed, take photographs and provide other documentation showing its significance in relation the food products.

Additional guidance can be found in 21 CFR as follows:
1. Part 110.20(b) - Plant Construction and Design,
2. Part 110.40(a) - Equipment and Utensils,
3. Part 110.35(c) - Pest Control,
4. Part 110.10(b) - Personnel Cleanliness.

Additional guidance can be found in 40 CFR Part 180 - Tolerances and Exemptions For Pesticides in Food Administered by The Environmental Protection Agency as follows:
1. Part 180.521 - Fumigants for grain-mill machinery; tolerances for residues, and
2. Part 180.522 - Fumigants for processed grains used in production of fermented malt beverages; tolerances for residues.

Be alert for:
1. Possible PCB contamination. Articles containing PCBs (e.g., transformers, PCB containers stored for disposal, electrical capacitors) must be marked with prescribed labeling to show they contain PCBs. No PCB-containing heat exchange fluids, hydraulic fluids or lubricants are allowed used in food plants. All PCB storage areas must be marked to show the presence of PCBs. Observe food plant transformers for possible leakage. If observed, determine if food items are stored in the area, and sample for PCB contamination. If PCBs are encountered in a food establishment, immediately advise management this is an objectionable condition and advise your supervisor.
2. Possible mix-up of pesticides or industrial chemicals with food raw materials.
3. Improperly stored pesticides or industrial chemicals (lids open, torn bags in close proximity to foods, signs of spillage on floors, pallets, shelves, etc.).
4. Incorrect application methods including excessive use. Many pesticide labels give instructions for use and precautions on the container.
5. Improper disposal or reuse of pesticide or industrial chemical containers.
6. Evidence of tracking powder or improper use of bait stations or baited traps.
7. Improper handling of equipment. Movable or motorized equipment used for handling possible chemical
5.4.7.1.4 - OTHER

use, product or yourself.
caution to prevent contamination of the immediate area of samples are to be collected to document misuse, exercise industrial chemical contamination noted or suspected. If Fully document the exact nature of any pesticide or use containers were used for food products.

If pesticide misuse is suspected, obtain the following information;
1. Name of exterminator and contract status,
2. Name of pesticide,
3. Name of pesticide manufacturer,
4. EPA registration number,
5. Active ingredients, and
6. Any significant markings on pesticide containers.

Fully document the exact nature of any pesticide or industrial chemical contamination noted or suspected. If samples are to be collected to document misuse, exercise caution to prevent contamination of the immediate area of use, product or yourself.

5.4.7.2 - Microbiological Concerns

Contamination of food products by bats, birds and/or other animals is possible in facilities where food and roosting facilities are available. Examine storage tanks, bins, and warehousing areas to determine condition and history of use. There have been instances where empty non-food use containers were used for food products.

5.4.7.2.1 - PROCESSING EQUIPMENT

Document the addition, or possible addition of pathogenic microorganisms from accumulated material due to poorly cleaned and/or sanitized processing equipment

Observe and report the firm's clean up procedures and the condition and cleanliness of food contact surfaces before production starts, between production runs and at the end of the day. Document any residue on food contact
surfaces of equipment, especially inside complex equipment not easily cleaned and sanitized. Report firms clean-up procedures in depth, since it may lend significance to insanitary conditions of residues on the plant machinery which are left to decompose overnight or between shifts. Where possible, observe equipment both before and after cleaning to assess it adequacy. Observations of residues on plant machinery can dramatically document the addition of pathogenic microorganisms, if present, into the product.

Identify any vectors of contamination (e.g. birds, rodents, insects, foot traffic, etc.), and describe sources and the routes of contamination from them to the product. Support this with your actual observations.

5.4.7.2.2 - EMPLOYEE PRACTICES

Document any poor employee practice and how they have or would provide a route for contaminating the product. For example, did employees (number/time of day) fail to wash and sanitize their hands at the beginning of processing, after breaks, meals, or after handling materials likely contaminated with a microbial pathogen, etc.; and then handle the finished product. Did employees handle product in an insanitary manner (cross contaminating raw product with cooked product, etc.; how many, how often).

5.4.7.3 - Storage

Evaluate the storage of finished products in the same manner as for raw materials. Determine if products are stored to minimize container abuse, facilitate proper rotation, and adherence to the storage requirements. This includes refrigeration temperatures, critical temperature tolerance, aging of products, and proper disposition of distressed stock.

5.4.7.3.1 - FOOD TRANSPORT VEHICLES

During food sanitation inspections, (See IOM 5.2.2.2 regarding issuance of FDA 482, Notice of Inspection while inspecting vehicles.), conduct inspections of food transport vehicles to include:

1. Evidence of insanitary conditions,
2. Conditions which might lead to food adulteration,
3. Physical defects in the vehicle,
4. Poor industry handling practices.

The following types of transport vehicles should be covered:

1. Railroad boxcars, both refrigerated and non-refrigerated, and hopper cars.
2. Any type of truck used to transport foods; both refrigerated and non-refrigerated.
3. Use extreme caution, if it is necessary to inspect tank railcars or tank trucks. Usually this coverage will be limited to determining what was transported in the tank previously and was the tank cleaned and/or sanitized as necessary between loads.

4. Vessels used to transport food in I/S commerce. Direct coverage primarily to intercoastal type vessels, including barges.

Coverage should be limited to food transport vehicles used for long haul (I/S) operations. Long haul vehicles are defined as those which travel at least 150 miles between loading and unloading or which do not return to the point of loading at the end of the day.

Regulatory actions are possible if unfit cars are loaded and, as a result of loading, adulteration occurs. Fully document any violations noted with appropriate samples and photographs. When vehicle insanitation is observed, it is imperative the carrier's and shipper's responsibility for the food adulteration be documented by appropriate evidence development, such as:

1. The nature and extent of the conditions or practices, and
2. The mechanical or construction defects associated with the food transport vehicle.
3. Individual responsibility for vehicle or trailer cleaning, vehicle assignments, load assignments, etc.

If gathering evidence about a single carrier, seek a series of occurrences at numerous locations involving as many different shippers as possible.

Basically two types of vehicles will be covered.

5.4.7.3.2 - VEHICLES AT RECEIVERS

When inspecting receivers of food products, examine the food transport vehicle prior to or during unloading. Make a preliminary assessment of food product condition, then inspect the vehicle after unloading to determine its condition and whether the unloaded food may have been contaminated during shipment. If the food appears to have been adulterated, collect a sample(s) for regulatory consideration. Samples collected from vehicles, which have moved the product in interstate commerce are official samples. You may also collect Documentary (DOC) Samples from the vehicle to substantiate the route of contamination.

5.4.7.3.3 - VEHICLES AT SHIPPERS

When inspecting shippers of food products, examine the food transport vehicle just prior to loading to determine its sanitary/structural conditions. If the vehicle has significant sanitation or structural deficiencies, notify the shipper of these conditions and of the possibility of product adulteration. If the shipper loads food aboard the vehicle, alert your supervisor so he/she can contact the FDA district where the consignee is located for possible follow-up. You may also collect samples from the load. These samples will become official when the Bill of Lading is issued.

5.4.8 - DISTRIBUTION
Report the general distribution pattern of the firm. Review interstate shipping records or invoices to report shipment of specific lots. If access to invoices or shipping records is not possible, observe shipping cartons, loading areas, order rooms, address stencils, railroad cars on sidings, etc., to determine customer names, addresses and destination of shipments. If no products are suspect, obtain a listing of the firm’s larger consignees.

5.4.8.1 - Promotion and Advertising

Determine the methods used to promote products and how the products reach the ultimate consumer. Determine what printed promotional materials are used and whether they accompany the products or are distributed under a separate promotional scheme. Check on the possibility of oral representations, i.e., door-to-door salesmen, spieler, etc. and obtain copies of brochures, pamphlets, tearsheets, instructions to salespersons, etc. Where indicated, obtain the lecture schedule of any promotional lecture program. If applicable, determine the general pattern of the media used for promotion and advertising.

5.4.8.2 - Recall Procedure

Determine the firm’s recall procedure. Audit enough records to determine the effectiveness of established procedures. Report if there is no recall procedure.

5.4.8.3 - Complaint Files

Review the firm’s complaint files. Where possible, copy the names and addresses of representative complainants; include a brief summary of each significant complaint in the EIR.

Identify who reviews complaints and their qualifications. Describe the criteria used by the firm in evaluating the significance of complaints and how they are investigated. Determine if records are kept of oral and telephone complaints. See IOM 5.2.8 for discussion of complaints with management and IOM 5.10.4.3.11 for reporting of complaints in the EIR.

Complaints may not be filed in one specific file, but may be scattered throughout various files under other subject titles including Product name; Customer name; Injured party name; Adjustment File; Customer Relations; Repair orders, etc.

During the inspection investigate all complaints contained on FDA-2516 and FDA-2516a forms in the firm’s district factory jacket. See IOM 5.2.8, 5.4.1.1 and 5.10.4.3.11.

5.4.9 - OTHER GOVERNMENT INSPECTION

See IOM 3.1 for general procedures on cooperating with other Federal, State, and local officials.

During Establishment Inspections determine the specific type of inspection service and inspecting units, which cover the firm, such as the name of the federal, state, county, or city health agency or department. Obtain the name and title of the inspectiveal official, and general method of operation.

5.4.9.1 - Federal

Do not inspect firms, or those portions of the plant, subject to compulsory, continuous inspection under USDA’s Meat Inspection Act, Poultry Products Inspection Act, or Egg Products Inspection Act, except on specific instructions from your supervisor or assignment document.

Ingredients or manufacturing processes common to both USDA and FDA regulated products should be inspected by FDA. See IOM 3.2.1.4 for FDA-USDA Agreements in specific areas.

Provide routine FDA coverage of such firms as breweries and wineries, which may be intermittently inspected on a compulsory basis by the U.S. Treasury Department, U.S. Public Health Service, or other agencies.

All products inspected under the voluntary inspection service of the Agriculture Marketing Service (AMS), USDA, and the National Marine Fisheries Service (NMFS), US Department of Commerce, are subject to FDA jurisdiction and are usually given routine coverage. However, formal written Agreements or Memoranda of Understanding between FDA and other agencies are often executed and may govern the agreeing agencies' operations on this type of inspected plants. When assigned this type of plant for inspection, always check to see if an Agreement or a Memorandum of Understanding exists between FDA and the agency involved to determine the obligations of both agencies. See IOM 3.1.2.1 and 3.2.

If you are assigned to cover a Federally Inspected plant which is under either compulsory or voluntary inspection, present your credentials and an FDA 482 Notice of Inspection to management and:
1. Identify yourself to the inspector(s) and invite him/her to accompany you on the inspection but do not insist on their participation.
2. At the conclusion of the inspection, offer to discuss your observations and provide the in-plant inspector with a copy of your Inspectional Observations (FDA 483).

5.4.9.2 - State and Local

State and local officials usually have extensive regulatory authority over firms in their area regardless of the interstate movement or origin of the food products involved. Joint FDA-State or local inspections are frequently conducted. These are usually arranged by district administrative or supervisory personnel. See IOM 3.1.2 and 3.3.

5.4.9.3 - Grade A Dairy Plant Inspections

If you are assigned to conduct an inspection or sample collection at a milk plant that is covered under the Grade A
Milk program, which has milk and milk products labeled and sold as Grade A, you should verify the need to complete the assignment with your supervisor and the Regional Milk Specialist. Grade A milk plants, milk, and milk products labeled as Grade A are inspected by state inspectors and check rated by FDA's Regional Milk Specialists and you should not inspect these Grade A milk and milk products. Milk plants in the Grade A Milk program and covered by the Interstate Milk Shippers (IMS) program are identified in the Interstate Milk Shippers List of Sanitation Compliance and Enforcement Ratings. This reference lists the specific milk plant and each milk and milk product covered under the IMS program. These Grade A milk and milk products are covered by a MOU between the FDA and the states, which places primary inspectional responsibility with the state.

There are situations where you will need to conduct an inspection in a Grade A milk plant and cover products they manufacture which do not carry the "Grade A" designation (such as juices). Fluid milk and milk products, cultured/acidified milk and milk products, eggnog, cream(s) sour cream, and yogurt are all considered Grade A and are required to be labeled as Grade A. The Grade A milk plant may also manufacture milk and milk products which are optional for the Grade A designation, depending upon the particular state. Cottage cheese is considered a Grade A optional milk product. If the state does not require the Grade A designation for cottage cheese, then the cottage cheese will not be included in the IMS listing of Grade A milk and milk products for that specific milk plant. Also, if the Grade A milk plant is manufacturing condensed or dried milk or milk products or condensed or dried whey or whey products, which are optionally labeled as Grade A, then those milk or milk products must be IMS listed and are covered under the Grade A Milk Program. Note: This same Grade A milk plant may also be manufacturing non-Grade A versions of these condensed/dried milk or milk products or condensed/dried whey or whey products.

5.4.10 - FOOD STANDARDS

The Federal Food, Drug, and Cosmetic Act requires the Secretary of Health and Human Services to promulgate reasonable definitions and Standards for food to promote honesty and fair dealing in the interest of consumers. When a Standard becomes effective, it establishes the common or usual name for the article, defines the article and fixes its standard of identity. It is then the official specification for the food. The food industry actively participates in the development of a Standard, and supplies much of the data upon which the regulation is based.

The Food Standards (FS) Inspection is made to obtain data for use, together with information from other sources in developing a Food Standard. Food Standard inspections are also made to determine a firm's compliance with food standards regulations, when manufacturing a standardized food.

5.4.10.1 - Food Establishment Inspection

Food Standard (FS) inspection assignments usually originate from CFSAN. When an inspection is planned for the purpose of collecting data to support a proposed food standard regulation, the district may elect to advise the firm, if the CFSAN has not already done so. If the firm selected does not choose to cooperate, it may be necessary to visit additional plants in order to obtain the desired information. Selection of additional firms should be done in consultation with the CFSAN.

Some firms often contend their entire process and formulas are "trade secrets". Attempt to persuade management the term "trade secret" should only be used to cover the process and/or quantitative-qualitative formulation which is truly unique to the firm. In instances where the firm is reluctant to release any of the information requested, point out FDA will, within the limits of the Freedom of Information Act, make every effort to preserve the confidentiality of the composition, make-up, and production levels of the product through the use of codes, which cannot be traced back to the firm. Include as much of the compositional and processing information as you can in the body of the report, without violating the firm's confidence.

5.4.10.2 - Food Inspection Report

FS EI's may be used as exhibits at public hearings and are subject to review by any interested party.

Three copies of the report are prepared. The original and one copy will be submitted to the CFSAN and one copy kept for the district file. Sign the original and duplicates of the first and last pages of each report sent to the Center.

Divide the report into three sections.

5.4.10.2.1 - ESTABLISHMENT INSPECTION RECORD (EI RECORD)

In order to relate the sections of the report to each other and to any assignments, and to assure any parts of the reports made public will not be identified as to the name of the firm or individuals therein, each district will set up a master list of numbers. One number will be assigned to each establishment covered, e.g., "BLT FS-3". For each FS Inspection place the assigned number next to the firm name on the EI Record. All other pages of the report shall be identified only by this number, the name of the commodity, and date. Example: "EI R Frozen Fish Sticks 10-3-87 BLT FS-3". This indicates a FS EI of frozen fish sticks conducted by Baltimore District on 10-3-87 in a plant designated as #3.

Where a producer may be reluctant to release any of the information requested, point out the FDA will, within the limits of the FOIA, make every effort to preserve the confidentiality of the composition, make-up, and production levels of his product through the use of codes, which cannot be traced back to the firm.
5.4.10.2.2 - BODY OF REPORT

Prepare the body of the report following the narrative outline as for any other food EIR except for the restrictions below.

The body of the FS report should also contain information in regard to the approximate annual value and volume as well as the percent of interstate business for each product covered. This is necessary because the coversheet, which contains this information, identifies the firm and will not be made public. Processes and the listing of raw materials used by the firm, which are not restricted by the term "trade secret" should be included. Any opinions, recommendations, or other information obtained or offered by individuals interviewed should be reported. Any suggestions made by individuals interviewed regarding what should be placed in the Standards for the products covered should be included. All individuals interviewed, firm name, etc. should have an identifying code assigned.

The body of the report should not include names and titles of individuals, (including USDA, USDI, or other inspectors), trade secret information, labeling, trade names, formulas, sample numbers, firm name or location of plant (other than by state or region), shipments, or other distribution information, legal status, or regulatory history. This information will be placed in the “Special Information” section of the report.

5.4.10.2.3 - SPECIAL INFORMATION SECTION

This is a separate attachment to the EIR which lists the names and titles of individuals (including other government inspectors) and firms with a reference code for each. The EIR should refer only to "Mr. A.,” "Mr. B.,” "Firm X,” "Firm Y”, etc. Do not use the firm or individual's actual initials in the body of the report. Include all information excluded from the body of the report and mount all labels obtained during the EI Labels may be quoted in the body of the report, but do not identify the firm. List the "Special Information Sheet" in the FACTS endorsement section as an enclosure.

Supplemental Reports - If, because of an additional visit or visits to the same firm on the same project, it is necessary to prepare another EIR, flag the report with the same number as assigned to the original report. For example, mark the EI Record "BLT FS-3 Supplemental Report", and the remaining pages, "EIR Frozen Fish Sticks 10-25-87 BLT FS-3 Supplemental Report."

5.4.10.3 - VIOLATIVE INSPECTIONS

When an inspection made in connection with the Food Standards project shows insanitary or other conditions which are not germane to the assignment or in the District's opinion suggests regulatory action, an appropriate narrative of the violative conditions should be prepared as a Regulatory Addendum.

5.4.11 - RE-INSPECTIONS CONDUCTED UNDER SECTION 743 OF THE FD&C ACT

The FD&C Act, as amended by the FDA Food Safety Modernization Act of 2011 (FSMA) (P.L. 111-353) authorizes FDA to collect fees to cover costs related to specific domestic and foreign food facility re-inspections. FDA announced our intent to collect these fees beginning in fiscal year 2012 via a Federal Register Notice issued on August 1, 2011 (FRN).

Section 743 of the FD&C Act authorizes FDA to assess and collect fees to capture 100% of the costs related to certain domestic and foreign food facility re-inspections. The fee for re-inspection is to cover re-inspection related costs when an initial inspection, initiated on or after October 1, 2011, has identified violations “materially related to food safety requirements” of the FD&C Act. The re-inspection must be conducted specifically to determine whether compliance has been achieved.

5.4.11.1 - Conducting Inspections for which Fees can be Assessed

To be eligible for a re-inspection for which fees can be assessed under section 743 of the FD&C Act:

1. a firm must have a previous inspection with a final classification of Official Action Indicated (OAI) initiated on or after October 1, 2011.

2. a firm must be evaluated to determine whether corrective actions have been implemented and effective and compliance has been achieved by the firm to FDA’s satisfaction.

3. a firm must be determined to be in situations where FDA has found the noncompliance was materially related to food safety requirements of the FD&C Act

These inspectional assignments should be generated by the District Compliance Branch as part of the OAI follow-up inspection process. The assignment will indicate that the inspection is a Reinspection for violations materially related to food safety requirements of the FD&C Act.

5.4.11.1.1 - REGULATED INDUSTRY NOTIFICATION

During every food and feed facility inspection, a copy of the Information Sheet - Assessment of Reinspection and Recall Fees by the FDA the FDA, should be provided to firm management at the same time the FDA-482, Notice of Inspection, is issued. Issuance of the Information Sheet should be documented as per section in the Administration Section of the Establishment Inspection Report (EIR). See IOM 5.10.4.3.3.

For foreign facility inspections, the Consumer Safety Officer (CSO) will provide copies of both the English version and the translated version, if available, during the
opening interview of the inspection. During a foreign facility inspection, the CSO shall identify and document the United States (US) Agent for that facility in the Administrative Data section of the EIR.

When Administrative and/or Enforcement actions are taken, standard language will be included in the official post-inspectional correspondence informing responsible firm management that subsequent re-inspections for the violations documented will be subject to fees under section 743 of the FD&C Act. When initiating a re-inspection, remind firm management of the fees that will be incurred for this inspection. Additionally, provide firm management with a copy of the Information Sheet – Assessment of Reinspection and Recall Fees by the FDA when issuing the FDA-482, Notice of Inspection and document the issuance in the EIR.

Firm officials can direct any billing questions to the “ORA FSMA User Fee Billing” email address (ORAFSMANeededFees@fda.hhs.gov).

5.4.11.1.2 - Fees for Non-compliance with a Recall Order

FDA will assess a fee for non-compliance with a recall order under Section 423(d) or 412(f) of the FD&C Act to cover food recall activities associated with such order. Non-compliance may include:

- not initiating a recall as ordered by FDA
- not conducting the recall in the manner specified by FDA in the recall order
- not providing FDA with requested information regarding the recall, as ordered by the FDA

Section 743(a)(1)(B) of the FD&C Act states that the fee is to be paid by the responsible party for domestic facilities and an importer who does not comply with a recall order under Section 423(d) or 412(f) of the FD&C Act. In other words, the paying the fee would be the party that received the recall order.

The fee is based on the number of direct hours spent on taking action in response to the firm’s failure to comply with a recall order. Types of billable activities could include:

- conducting recall audit checks
- reviewing periodic status reports
- analyzing status reports and audit check results
- conducting inspections
- travel to and from locations
- continuing monitoring of product disposition

Supervisory review and administrative support time does not need to be tracked and reported on the timesheet as those activities have already been accounted for in the hourly rate. Record each event activity description, amount of time per activity and any relevant comments.

5.4.11.1.3 - RE-INSPECTION ASSIGNMENT GENERATOR

The District Compliance Branch will generate OAI follow-up inspection assignments in FACTS. The re-inspection designation should be noted in the FACTS assignment subject line. The re-inspection designation for foreign inspections will be accomplished by DFFPOI or DMPTI upon communication from the respective Center. In the Background section of the FACTS assignment, include an affirmation that the violations are materially related to a food safety requirement of the FD&C Act.

5.4.11.1.4 - RE-INSPECTION REPORTING

When assigned a re-inspection for which FDA can assess fees under section 743 of the FD&C Act, the lead CSO will be responsible for obtaining the FACTS Assignment and the FSMA Fee Re-Inspection Time Reporting for Inspection Hours sheets. The FACTS Assignment Sheet must be obtained prior to setting the FACTS inspection assignment to Complete. Each CSO involved in preparing, conducting, and reporting a FSMA fee re-inspection must accurately complete, sign and date the FSMA User Fee Re-inspection Timesheet.

Districts should ensure that any and all reinspection time is properly tracked to allow for accurate and prompt billing by the agency. Both the FSMA Fee Re-Inspection Time Reporting for Inspection Hours and FSMA Reinspection Cover Sheet forms are designed for efficient tracking of time and activities. On a daily basis, Investigators will record the date, participant(s) name, activity description, amount of time per activity and any relevant comments. Please note that all time expended by the Investigator is reimbursable including time spent for:

1. Preparation
2. travel to and from activities (including travel for road trip
3. and foreign inspections)
4. conducting the inspection
5. collecting samples
6. conducting field examinations
7. report writing

When reporting time into the FACTS database, the “Reimbursable” check box must be checked on the Inspection Accomplishment Hours screen. Exhibit 5-18 shows where this box is located.

When a supervisor reviews the EIR and records the FACTS endorsement and initial classification in the FACTS EI Record, they must confirm the Reimbursable
check box is checked in FACTS. The supervisor will complete the FSMA Re-Inspection Fee Cover Sheet and submit this document with the FSMA Fee Re-Inspection Time Reporting for Inspection Hours sheet and FACTS Assignment Sheet completed by the investigator. The supervisor should ensure separation of any time spent on inspectional issues not materially related to a food safety requirement of the FD&C Act. After review, the supervisor will sign and date the FSMA Fee Re-Inspection Time Reporting for Inspection Hours sheet and coversheet. For a foreign re-inspection, the supervisor should ensure that the US Agent information is captured in the FSMA Re-Inspection Fee Cover Sheet and in the Administrative Data section of the EIR. The User Fee Re-Inspection package should be provided to the District Compliance Branch and contain:

1. FSMA Re-Inspection Fee Cover Sheet
2. FACTS Assignment Sheet
3. FSMA Fee Re-Inspection Time Reporting for Inspection Hours sheet
4. Completed EIR with attachments and exhibits

The District Compliance Branch shall submit to the Office of Resource Management (ORM) at “ORA FSMA User Fee Billing” (ORAFSMAUserFeeBilling@fda.hhs.gov):

1. FSMA Re-Inspection Cover Sheet
2. FACTS assignment sheet
3. FSMA User Fee Re-Inspection Time Reporting sheet

ORA/OO/OFFO/DFFPOI has added a web page to their intranet site for resources and guidance related to FSMA (including fee, FA and Investigator form links/ information) which may be accessed at: http://inside.fda.gov:9003/ORA/OfficeofRegionalOperation s/DivisionofFieldInvestigations/ucm273532.htm.

Districts should consult with CFSAN- Office of Compliance, Division of Enforcement (HFS-605), CVM- Office of Compliance and Surveillance, Division of Compliance (HFV-230), or the Office of Chief Counsel, as appropriate, if questions arise in determining whether a non-compliance is materially related to a food safety requirement of the FD&C Act.

Office of Resource Management (ORM) has established a FSMA Fee Team. This team will be responsible for receiving billing packages from the field and processing them for billing by the Office of Financial Management (OFM). ORM will also run reports using ORADSS to identify inspections checked as “Reimbursable” and reconcile with billing packages received. The FSMA Fee Team will also field questions from industry regarding billing using the “ORA FSMA User Fee Billing” email address at ORAFSMAUserFeeBilling@fda.hhs.gov.

When samples are collected for analysis and compliance purposes, District Compliance Branches will receive Collection Reports (CRs)/Analyst Worksheet packages for review as part of the regulatory process. District Compliance Branches will also be responsible for reviewing Lab Accomplishment Hours in FACTS and entering those hours on both the FSMA Fee Re-Inspection Time Reporting for Inspection Hours and FSMA Reinspection Cover Sheet forms. Please note that Analyst time is reported in FACTS separately from any inspectional activities. To review Lab Accomplishment Hours in FACTS: Navigate to the Sample Summary for the analysis, click on Options and select Lab Accomplishment Hours.

5.4.11.1.5 – RE-INSPECTION COMPLIANCE BRANCH ACTIONS

For domestic inspections, District Compliance Branches, along with OE/CFSAN/CVM, will determine if violations are “materially related to a food safety requirement of the FD&C Act.” For foreign inspections, CFSAN or CVM will make this determination. Once it is determined the violations are related, the reinspection of the facility, specifically for compliance follow-up purposes, will be subject to fees.

As per section 4-1-10 Warning Letter Format of the Regulatory Procedures Manual (RPM), for Warning Letters (WL) based on inspections of food facilities classified as OAI that identify noncompliance(s) material to a food safety requirement of the Act, District Compliance Branches and Center Offices of Compliance must include the following statement for domestic or foreign facilities, as applicable. [Bold type in brackets] indicates that appropriate language must be inserted: “Section 743 of the Act (21 U.S.C. 379j-31) authorizes FDA to assess and collect fees to cover any reinspection-related costs.”

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5.4.12 - PESTICIDES

5.4.12.1 - Pesticide Inspections

The objective of a Pesticide Inspection is to determine the likelihood of excessive residues of significant pesticides in or on products in consumer channels, and to develop
sources of information for uncovering improper use of pesticide chemicals.

This requires directing coverage to two major areas:
1. Pesticide practices in the production and processing of field crops.
2. Application of pesticide chemicals in establishments storing and processing raw agricultural products.

Pesticide coverage must be provided during all food establishment inspections. Coverage of raw agricultural products will generally be on a growing-area basis.

Problem areas include:
1. Improper use of pesticides around animals - gross misuse of sprays and dips in animal husbandry may result in pesticide residues in foods.
2. Use of contaminated animal feeds - waste and spent materials from processing operations may contain heavy concentrations of pesticide residues, which were present in the original commodity. See Compliance Policy Guide 575.100.
3. Past pesticide usage - past pesticide practices on growing fields. Past use of persistent pesticides may result in excessive residues in the current food crop. You may need to check on pesticide usage for several years prior to an incident to ensure you gather enough information. Some pesticides last for many years in the environment.

5.4.12.2 - Current Practices

Cooperative Activities - important sources of information relative to evaluating the "Pesticide Environment" include:
1. At the start of the growing season, spray schedules recommended for each crop by county agents, state experiment stations, large pesticide dealers, farmers cooperatives, et al should be obtained.
2. Visits to agricultural advisors may provide information relative to heavy infestation of insect pests and fungal infections on specific crops in specific areas.
3. Daily radio broadcasts in most agricultural areas may provide information on spray schedules, insect pests, harvesting and shipping locations, etc.
4. Field employees of fruit and vegetable canning and freezing plants usually recommend spray schedules, pesticides, and harvesting schedules for products produced by contract growers.
5. United States Weather Bureau Offices and their reports will provide data on weather conditions, which may effect insect growth and their development, size of fruit or leaf growth, and dissipation of pesticide chemicals.
6. USDA Market News Service daily price quotations, and weekly quotations in trade magazines provide information regarding harvesting schedules since market prices are indicators of how quickly a crop will be harvested in a given area. Growers who have the opportunity to obtain high prices may harvest their crops without regard to recommended pre-harvest intervals.

7. State Colleges of Agriculture seminars or short courses on food and vegetable production may alert you to significant departures from usual agricultural practices. Prior approval to attend such meetings should be secured from your supervisor.
8. Pesticide suppliers and distributors may provide information on spray practices, schedules, and the name and address of growers, etc.

NOTE: The U.S. Department of Agriculture has a Pesticide Data Program (PDP), which provides data on pesticide use and residue detection. This program helps form the basis for conducting realistic dietary risk assessments and evaluating pesticide tolerances. Coordination of this program is multi-departmental, involving USDA, EPA and FDA, covered by a MOU (Federal Cooperative Agreements Manual). As a part of this program USDA collects data on agricultural chemical usage, and factors influencing chemical use, and collects pesticide residue data through cooperation with nine participating states. USDA provides this data to EPA, FDA and the public. Several USDA publications are listed below as reference material.

The contact point at USDA for pesticide residue matters is:
Martha Lamont, Director
Monitoring Program Office, Science Division
Agricultural Marketing Service, USDA
8609 Sudley Road, Suite 206
Manassas, VA 20110
703-330-2300

Reference materials - the following reference materials provide background and data necessary or helpful in evaluating current practices. This material should be available at the District office.
1. Pesticide Chemicals - Regulations under the Federal Food, Drug and Cosmetic Act on tolerances for pesticides in food administered by the Environmental Protection Agency (EPA). (See 40 CFR 185)
2. EPA's Pesticide Regulations - Tolerances for Raw Agriculture Products. (See 40 CFR 180)
3. EPA's Rebuttable Presumption Against Registration (RPAR) List.
4. Pesticide Index. - By William J. Wiswesser. A publication containing information on trade names, composition and uses of commercial pesticide formulations.
5. The Daily Summary or Weekly Summary. News releases and reports from USDA.
8. Annual Pesticide Data Summary
9. Reports from USDA's Crop Reporting Board.
10. USDA's Pesticide Assessment Reports.

5.4.12.3 - Growers
Preliminary investigation of growing areas at the start of the season will provide data necessary for district work planning including production schedules, types and acreage of crops, pesticides used and the names and addresses of growers and shippers.

Growing Dates - The significant growing dates relative to pesticide usage are as follows:
1. Planting date,
2. Date of full bloom, and
3. Date of edible parts formation.

Harvest Dates - The dates of the anticipated harvest season will provide planning information relative to pre-harvest application and shipping.

Acreage - This will provide volume information for work planning.

5.4.12.3.1 - PESTICIDE APPLICATION

Ascertain the actual pesticide application pattern for each crop. Look for objective evidence to document actual grower practice. Check the grower's supply of pesticide chemicals, look for used pesticide containers, visit his source of supply, etc. Check spraying and dusting practices. Establish if pesticide chemicals are used in such a manner that excessive residues might result.

The following information provides a basis for evaluating pesticide usage:
1. Pesticide Chemical Applied - List the common name if there is no doubt as to the chemical identity of the pesticide. Include labeling indications and instructions.
2. Method of Application - Describe the method of application i.e., ground rig, airplane, greenhouse aerosol, hand, etc.
3. Formulation - Describe the formulation i.e., wettable powder, emulsifiable concentrate, dust, granules, aerosol, etc. Express as pounds of active ingredient per gallon or percent wettable powder.
4. Number of Applications and Dates.
5. Rate of Last Application - Calculate the amount of active ingredient per acre.
6. Pre-Harvest Interval (PHI) - Calculate the number of days between the day of the last application of pesticide and the harvest date or anticipated harvest date. Compare to the PHI.
7. Visible residue on grower's crop.
8. Summary of Usage - Determine the USDA Summary Limitations and evaluate the responsible usage.

5.4.12.3.2 - PESTICIDE MISUSE/DRIFT/SOIL CONTAMINATION

Pesticide residues, which exceed established tolerances, action levels, or "regulatory analytical limits", may be caused by pesticide misuse which can include:
1. Excessive application of a chemical on a permitted crop.
2. Failure to follow labeled time intervals between the last pesticide application and harvest.

3. Use of a non-approved pesticide on a crop.
4. Failure to wash a crop when pesticide labeling requires it (e.g., for certain EBDC's).

Other conditions, which may cause illegal residues, include spray drift and soil contamination.

Drift may be documented by determining which crops and pesticides have been grown/used in fields adjacent to those sampled. Determine direction of prevailing winds and wind condition on the day of spraying. Selective sampling will aid in determining if drift occurred. Compliance Samples collected to document pesticide drift should be Flagged as a Pesticide Sample and noted in the Remarks section of the CR as "Drift Sample - Maintain as Individual Subs".

Soil contamination by compounds, which are relatively stable in the environment, may cause systemic uptake of the compounds by growing crops. Follow-up investigations to violative samples may, in some limited cases, include soil samples as an attempt to determine the source of the contaminant. Do not routinely collect soil samples.

5.4.12.4 - Packers and Shippers

Follow the same general procedure as in IOM 5.4.12.3. Observe and report the following:
1. Treatment Before Shipping - This may include stripping of leaves, washing, vacuum cooling, application of post-harvest preservative chemicals, use of cartons with mold-inhibiting chemicals, waxes, colors, fumigation, etc.
2. Identification of Growers' Lots - Determine procedure or methods used to maintain the identity of each grower's lot. Provide the code and key if any.
3. Labeling - Quote labeling or brand names.
4. Responsibility - Determine whether the packer or shipper knows what sprays have been used on the products shipped.

5.4.12.5 - Pesticide Suppliers

Pesticide suppliers should be visited routinely during growing-area coverage. They may provide valuable information about pesticides being used on various crops in the growing area. Some suppliers may suggest spray schedules or advise growers about pesticide usage.

Determine what representations were made by the manufacturer of pesticide chemicals for which there is only a temporary tolerance or experimental permit. Get copies of any correspondence relating to sale and use of these products. Obtain names of growers to whom sales are made if such sale was not for use on acreage assigned under the experimental permit. Collect Official Samples of any crops treated with the pesticide.

5.4.12.6 - Pesticide Applicators

Pesticide applicators may provide valuable information about pesticides being used on various crops in the
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growing area. Interview several pesticide applicators, particularly those using airborne equipment. Determine the pesticide chemicals, their formulation, and on what crops they are currently being applied. Determine who supplies the pesticides and how they are prepared to assure proper concentration. If state law requires the applicator to keep a record of each spray application, request permission to review such records. Determine what steps are taken to assure drift on adjoining crops does not result in violative residues. Where there is likelihood of drift, collect Selective Samples from adjoining fields.

5.4.12.7 - Sample Collections

See IOM Sample Schedule Chart 3 - Pesticides.

SUBCHAPTER 5.5 - DRUGS

5.5.1 - DRUG INSPECTIONS

Authority for inspection is discussed in IOM 2.2. FD&C Act Sections 501(a)-(d) and 501(j) [21 U.S.C. 351(a)-(d), (j)] describe the ways in which a drug may be or may become adulterated. Section 502 of the FD&C Act [21 U.S.C. 352] does the same, with respect to misbranding. Section 505 of the FD&C Act [21 U.S.C. 355] requires that new drugs be approved by FDA. Therefore, the purposes of a drug inspection are:

1. To evaluate a firm's adherence to the concepts of sanitation and good manufacturing practice; i.e., production and control procedures include all reasonable precautions to ensure the identity, strength, quality, and purity of the finished products;
2. To identify deficiencies that could lead to the manufacturing and distribution of products in violation of the Act, e.g., non-conformance with Official Compendia, super/sub potency, substitution;
3. To determine whether a firm is distributing drugs that lack required FDA approval including counterfeit or diverted drugs;
4. To obtain correction of those deficiencies;
5. To determine if new drugs are manufactured by the same procedures and formulations as specified in the New Drug Application documents;
6. To determine the drug labeling and promotional practices of the firm;
7. To ensure the firm is reporting NDA field alerts as required by 21 CFR 314.81 and Biological Product Deviation Reports (BPDRs) for therapeutic biological products as required by 21 CFR 600.14;
8. To determine if the firm is complying with the requirements of the Prescription Drug Marketing Act (PDMA) and regulations; and
9. To determine the disposition of Drug Quality Reports (DQRs) received from the Drug Surveillance and Data Reporting Branch (DSDRB)/CDER; and
10. To determine if the firm is complying with postmarket Adverse Drug Experience reporting requirements as required by 21 CFR sections 310.305 (prescription drugs without approved NDA/ANDA), 314.80, 314.98, and 314.540 (application drug products), and 600.80 (therapeutic biological products), and Section 760 of the FD&C Act (non-application nonprescription products) [21 U.S.C. 379aa].

5.5.1.1 - Preparation and References

Become familiar with current programs related to drugs. Determine the nature of the assignment, i.e., a specific drug problem or a routine inspection, and if necessary, consult other district personnel, such as chemists, microbiologists, etc., or center personnel, such as office of compliance staff. Review the district files of the firm to be inspected including:
1. Establishment Inspection Reports,
2. District Profiles,
3. OTC monographs and other pertinent references for non-application products,
4. Drug Applications (New, Abbreviated and Investigational) and the Knowledge Transfer Memo,, if the Center has provided it for a specific pre-approval inspection,
5. Therapeutic Biologics License Applications,
6. Sample results,
7. Complaints and Recalls,
8. Regulatory files,
9. Drug Quality Reports (DQRs), NDA Field Alert Reports (FARs), and Biological Product Deviation Reports (BPDRs),
10. Drug Registration and Listing

During this review identify products which:
1. Are difficult to manufacture,
2. Are complex dosage forms,
3. Require special tests or assays, or cannot be assayed,
4. Require special processes or equipment,
5. Are new drugs and/or potent low dosage drugs,
6. Are misbranded, unapproved, fraudulent, or compounded drugs containing ingredients that have been withdrawn or removed from the market for safety or effectiveness reasons, or compounded drugs that contain bulk active ingredients that are not components of FDA-approved drugs.

Review the factory jacket, FACTS OEI and registration/listing data, and all complaint reports which are marked follow-up next inspection. These complaints are to be investigated during the inspection and discussed with management. See IOM 5.2.7.

Become familiar with current regulations and programs relating to drugs, CP 7356.002, et al. When making GMP inspections, discuss with your supervisor the advisability of using a microbiologist, analyst, engineer, or other technical personnel to aid in evaluating those areas of the firm germane to their expertise. Review the FD&C Act, Chapter V, Drugs and Devices, Review parts of 21 CFR 210/211 applicable to the inspection involved and Bioavailability (21 CFR 320). In the case of APIs, review FD&C Act section 501(a)(2)(B) [21 U.S.C 351(a)(2)(B)]
and the ICH industry guideline entitled "Q7A Good Manufacturing Practice Guidance for Active Pharmaceutical Ingredients."

Review the current editions of the United States Pharmacopeia (USP), and Remington's Pharmaceutical Sciences for information on specific products or dosage forms. For compounding pharmacies, review USP Chapters 795 and 797. See IOM 1.10.3 for special regulatory information by product category.


Before conducting drug preapproval inspections (CP 7346.832) it is important to be familiar with the application and coordinate accomplishment of Center goals communicated by (1) inspecional memos, (2) pre-inspection briefings, and/or (3) Center participation on the inspection team.

Before conducting an inspection that may involve postmarketing adverse drug experience reporting, you should review 21 CFR Sections 310.305, 314.80, 314.98, 314.540, and 600.80, Section 760 of the FD&C Act [21 U.S.C. 379aa], CP 7353.001, and the training video, 'Field Investigators: Adverse Drug Effects (ADE) Detectives.'

The Office of Manufacturing and Product Quality (OMPQ) in CDER has established two mechanisms for you to obtain technical assistance before, during, or after an inspection:

1. Office of Manufacturing and Product Quality (OMPQ) Subject Contacts. This list contains the names and phone numbers of OMPQ individuals identified as technical specialists in various areas.

2. Questions and Answers on Current Good Manufacturing Practices for Drugs. This forum is intended to provide timely answers to questions about the meaning and application of CGMPs for human, animal, and biological drugs, and to share these widely. These questions and answers generally clarify statements of existing requirements or policy.

5.5.1.2 - Inspecional Approach

Follow Compliance Program Guidance Manual (CP) 7356.002 and others as appropriate when conducting CGMP inspections. In-depth inspection of all manufacturing and control operations is usually not feasible or practical. A risk-based systems audit approach is recommended in which higher risk, therapeutically significant, medically necessary and difficult to manufacture drugs are covered in greater detail during an inspection. (Note: The status of a drug as medically necessary is determined by CDER. For information contact Office of Compliance/Recalls and Shortages Branch via email at cder recalls@fda.hhs.gov) The latter group includes, but is not limited to, time release and low dose products, metered dose aerosols, aseptically processed drugs, and formulations with components that are not freely soluble.

CP 7356.002 incorporates the systems-based approach to conducting an inspection and identifies six (6) systems in a drug establishment for inspection: Quality, Facilities and Equipment, Materials, Production, Packaging and Labeling and Laboratory Control Systems. The full inspection option includes coverage of at least four (4) of the systems; the abbreviated inspection option covers of at least two (2) systems. In both cases, CP 7356.002, indicates the Quality System be selected as one of the systems being covered. During the evaluation of the Quality System it is important to determine if top management makes science-based decisions and acts promptly to identify, investigate, correct, and prevent manufacturing problems likely to, or have led to, product quality problems.

When inspecting drug manufacturers marketing a number of drugs meeting the risk criteria, the following may help you identify suspect products:

1. Reviewing the firm's complaint files early in the inspection to determine relative numbers of complaints per product.

2. Inspecting the quarantine, returned, reprocessed, and/or rejected product storage areas to identify rejected product.

3. Identifying those products which have process control problems and batch rejections via review of processing trends and examining reviews performed under 21 CFR 211.180(e).

4. Reviewing summaries of laboratory data (e.g., laboratory workbooks), OOS investigations, and laboratory deviation reports.

5.5.2 - DRUG REGISTRATION & LISTING

Registration and listing is required whether or not interstate commerce is involved. See Exhibit 5-12 and IOM 2.9.1.1 for additional information.

Two or more companies occupying the same premises and having interlocking management are considered one establishment and usually will be assigned a single registration number. See IOM 5.1.1.11 - Multiple Occupancy Inspections for additional information.

Independent laboratories providing analytical or other laboratory control services on commercially marketed drugs must register.

FACTS will indicate if the establishment is registered for the current year. If you determine registration and listing is required, advise your supervisor. After checking for past registration, cancellation, etc., the district will provide the firm with the proper forms and instructions.

Each establishment is required to list with FDA every drug in commercial distribution, whether or not the output of such establishment or any particular drug so listed enters
interstate commerce. During the establishment inspection, you should remind the firm of its responsibilities for ensuring its drug listing accurately reflects the current product line and updating its listing as necessary to include all product changes, NDC changes, and discontinuations in accordance with 21 CFR 207. If registration and listing deficiencies are found, document it in your EIR, collect a documentary sample and/or contact your supervisor.

5.5.3 - PROMOTION AND ADVERTISING

21 CFR 202.1 which pertains only to prescription drugs, covers advertisements in published journals, magazines, other periodicals, and newspapers, and advertisements broadcast through media such as radio, television, and telephone communication systems. Determine what department or individual is responsible for promotion and advertising and how this responsibility is demonstrated. Ascertain what media (radio, television, newspapers, trade journals, etc.) are utilized to promote products.

Do not routinely collect examples of current advertising. Advertising should be collected only on assignment, or if, in your opinion, it is clearly in violation of Section 502(n) of the FD&C Act [21 U.S.C. 352 (n)] or 21 CFR 202.1.

5.5.4 - GUARANTEES AND LABELING AGREEMENTS

Determine the firm's policies relative to receiving guarantees for raw materials, and issuing guarantees on their products. Also determine firm's practices regarding shipment of unlabeled drugs under labeling agreements. See IOM 5.3.7.2.

5.5.5 - OTHER INSPECTIONAL ISSUES

5.5.5.1 - Intended Use

Please see the discussion of jurisdiction in section IOM 5.10.4.3.6.

5.5.5.2 - Drug Approval Status

The investigator should ascertain whether the drugs manufactured by the firm are covered by an NDA, ANDA, OTC monograph, or marketed under a claim of DESI or "grandfather" status.

5.5.5.3 – Drug Status Questions

If you have questions about misbranding, new drug status, API/finished drug product status, drug/cosmetic, drug/food (dietary supplement) status, or compounded drugs, contact the Office of Unapproved Drugs and Labeling Compliance in CDER's Office of Compliance at 301-796-3100 or CDEROUNDLCMTRACK@CDER.FDA.GOV. In order to make these determinations, drug product labeling is needed.

In rare cases, a drug may be unapproved and inappropriate for marketing under any circumstances (i.e., it cannot be reconditioned or reformulated into a product appropriate for marketing). If you encounter products in this category, contact your supervisor to determine if a CGMP inspection is warranted.

5.5.5.4 - Drug/Dietary Supplement Status

In instances where the drug/dietary supplement status of a product is unclear, the investigator should collect all related labeling and promotional materials including pertinent Internet web sites. This labeling and promotional material is often useful in determining the intended use of a product (See 21 CFR 201.128). Labeling, promotional materials and Internet web sites often contain information, for example, disease claims, that can be used to determine the intended use of a product and thereby if it is a dietary supplement or a drug and an unapproved new drug.

5.5.5.5 - Approved Drugs

Check the current programs in your CP, Section 505 of the FD&C Act [21 U.S.C. 355] and 21 CFR part 314 for required information. You may take the district's copy of the NDA into the plant as a reference during the inspection. Document and report all deviations from representations in the NDA even though they may appear to be minor.

5.5.5.6 - Investigational Drugs

Follow the instructions in pertinent programs in your CP or as indicated in the specific assignment received.

5.5.5.7 - Clinical Investigators and/or Clinical Pharmacologists

Inspections in this area will be on specific assignment previously cleared by the Administration. Follow guidance in the CP or assignment.

5.5.5.8 – Delaying, Denying, Limiting or Refusing Drug Inspections

Use reasonable discretion when discerning whether action taken by a drug firm during an inspection constitutes delaying, denying, limiting, or refusing drug inspection. If you are unsure whether an action taken by a firm constitutes delaying, denying, limiting, or refusing drug inspection, contact your supervisor.

As needed, refer to the Guidance for Industry – Circumstances that Constitute Delaying, Denying, Limiting, or Refusing a Drug Inspection, for examples of firm actions that may cause a drug to be deemed adulterated under FD&C Act section 501(j). Remember, however, that these examples are not exhaustive, and that
guidance documents do not establish legally enforceable rights or responsibilities and are not legally binding on the firm or the agency. See IOM 1.10.1.

5.5.6 - CDMR BIO-RESEARCH MONITORING

Inspectional activities in the bio-research monitoring (BIMO) programs involve all product areas and Centers, including In Vivo Bio-equivalence, Good Laboratory Practice (GLP) for Non-Clinical Laboratories, Institutional Review Boards (IRB), Sponsors, Monitors, Contract Research Organizations, and Clinical Investigators (CI). In most instances, inspections conducted under this program will be done on assignment from the respective Center and occasionally with the participation of Center personnel as part of the inspection team.

During team inspections with Center personnel, the Field Investigator is the team leader. See IOM 5.1.2.5. The Compliance Program Guidance Manual (CP) for each program provides a description of the program and detailed instruction for conducting inspections.

Districts will make the initial classification of inspections and the Center issuing the assignment will make the final decision after review.

5.5.7 - ADVERSE EVENT REPORTING/Risk Evaluation and Mitigation Strategies (REMS)

FD&C Act section 760 [21 U.S.C. 379aa] and 21 CFR sections 310.305, 314.80, 314.98, and 314.540 require reporting of adverse events associated with the use of human drug products and section 600.80 requires reporting of adverse events associated with the use of biological products (including therapeutic biological products). Responsible firms include holders of NDAs and ANDAs, and manufacturers, packers and distributors that are named on the labels of all FDA approved drug products and all prescription drug products. Both foreign and domestic firms are required to develop written procedures and to maintain records related to adverse events. Firms must evaluate adverse event data to determine if the event has had a serious outcome such as death, disability, hospitalization, or was a life threatening event, and if the event was expected (labeled) or unexpected (unlabeled) for the product. Responsible firms must submit adverse event information to FDA in expedited or periodic reports, as described in the regulations.

Refer to the Compliance Program Guidance Manual (CP) (section 7353.001) for the description of the program and for detailed instructions for conducting inspections.

FD&C Act Section 505-1 [21 U.S.C. 355-1] gives the FDA the authority to require Risk Evaluation and Mitigation Strategies (REMS) for certain drugs to ensure that the benefits outweigh the risks.

Since every REMS program varies, the detailed instructions for conducting inspections will be given to the investigator prior to each inspection.

5.5.8 - DRUG INSPECTION REPORT

See IOM 1.1 English language requirement. The requirements in IOM 5.10.4.3, and any applicable Compliance Program Guidance Manuals can be used to help you prepare your report.

This does not cover the reporting requirements for a directed inspection with a narrow focus, such as a complaint follow-up or investigation into a recall. In those cases, use your judgment and guidance in IOM 5.10.4 about the depth of reporting required. Follow the instructions and format for a human drug inspection report as contained in IOM 5.10.4.2 and 5.10.4.3.

This human drug inspection report does not require full and detailed narratives for every area for every inspection. The firm's state of compliance, the previous inspectional report and information, complexity of operations and other aspects all are determinants in how much reporting will be necessary. In many cases, brief summaries addressing the format areas will be sufficient.

SUBCHAPTER 5.6 - DEVICES

5.6.1 - DEVICE INSPECTIONS

See IOM 2.2 for discussion of statutory authority.

The term "device" is defined in Sec. 201(h) of the FD&C Act [21 U.S.C. 321 (h)]. In-vitro diagnostics (21 CFR 809) are devices, as defined in 201(h) of the Act [21 U.S.C. 321 (h)], and may also be biological products subject to Section 351 of the PHS Act.

Inspections involving devices should be made only by those individuals qualified by training and experience in the device area. Electronic product radiation is defined in 21 CFR 1000. Because of the specific nature of inspections and investigations involving radiation, only personnel who have special training in this field should be assigned such work. However, others may participate for training purposes. Specific Compliance Program Guidance Manuals designate the type of individual and special training required for work in these areas.

CAUTION: Radiation-emitting devices and substances present a unique hazard and risk potential. Every effort should be taken to prevent any undue exposure or contamination. Monitoring devices must be used whenever radiation exposure is possible. Investigators should also be on the alert for, and avoid contact with, manufacturing materials and hazards associated with the manufacturing of many types of devices, which may present a threat to health, e.g., ethylene oxide, high voltage, pathogenic biomaterials, etc. See IOM 1.5 for additional safety information.
5.6.1.1 - Technical Assistance

Each region and some districts have engineers and radiological health personnel available for technical assistance and consultation. Do not hesitate to make use of their services.

Engineers, quality assurance specialists, and expert investigators in ORA/OO/OMPTO/Division of Medical Products and Tobacco Program Operations (DMPTPO), 301-796-0358, are available for on-site consultation and assistance in problem areas. The division's subject matter experts are also available by telephone for consultation and to answer questions regarding regulation and program interpretation and QS/GMP application. Additionally, the CDRH Office of Compliance enforcement divisions (organized by device product) can be contacted as necessary.

WEAC has various personnel (biomedical, sterility, electronic, materials, mechanical, nuclear and plastics engineers) available for telephone consultation and on-site assistance at 781-756-9700.

5.6.1.2 - Sample Collection During Inspection

Because of the limited funds available for samples and the relatively high cost of device samples, it is essential you consider, in consultation with your supervisor, the following factors before collecting a physical sample of a device:

1. If follow-up to a QS/GMP deviation, will sampling demonstrate the deviation and/or a defective product? Documentary Samples may be more suitable for QS/GMP purposes.
2. Likelihood of the analysis showing the device is unfit for its intended use.
3. Samples costing over $250.00.
4. Laboratory capability to analyze the sample. See IOM 4.5.5.3.6 for sample routing information.

If you are still uncertain, discuss with your supervisor and contact the CDRH Laboratory or WEAC 781-756-9700 for assistance.

Contact CDRH for assistance as follows:

In-vitro Diagnostic Devices - Office of Science and Technology (HFZ-113).

NOTE: Device samples do not require 702(b) portions. Include in the FDA 525 and with the C/R, if destined for different locations, a copy of the firm's finished device specifications, test methods and acceptance and/or rejection criteria.

5.6.1.3 - Types of Inspections

General device inspections will be conducted under various Compliance Programs found in the Compliance Program Guidance Manual. The majority of these will be QS/GMP inspections, but often the reason for the inspection will vary. For example, inspections may be conducted to assist the pre-market clearance process (PMA or Class III 510(k)), to specifically address MDR concerns, or to assure in-depth coverage of an aspect of manufacturing (sterility). The following describes some of these inspections.

5.6.1.4 - CDRH Bio-research Monitoring

Bio-research monitoring (BIMO) assignments for medical devices will generally be issued by the Center for Devices and Radiological Health (CDRH) (see IOM 5.5.6).

5.6.2 - MEDICAL DEVICE QUALITY SYSTEM/GOOD MANUFACTURING PRACTICES

Section 520(f) of the FD&C Act [21 U.S.C. 360j(f)] provides the Agency with authority to prescribe regulations requiring that the methods used in, and the facilities and controls used for, the manufacture, packing, storage, and installation of medical devices conform to good manufacturing practices. The medical device Quality System/Good Manufacturing Practices Regulation (QS/GMP) [21 CFR 820] became effective on June 1, 1997.

21 CFR 820 is established and promulgated under the authority of Sections 501, 502, 510, 513, 514, 515, 518, 519, 520, 522, 701, 704, 801 and 803 of the FD&C Act (21 U.S.C. 351, 352, 360, 360c, 360d, 360e, 360h, 360i, 360j, 360l, 371, 374, 381 and 383). Failure to comply with the provisions of 21 CFR 820 renders a device adulterated under Section 501(h) of the FD&C Act [21 U.S.C. 351(h)].

The regulations promulgated under 21 CFR 820 establish minimum requirements applicable to finished devices, as defined in 820.1(a). This regulation is not intended to apply to manufacturers of components or parts of finished devices, but instead recommended to them as a guide. In some special cases, components have been classified as finished devices (dental resins, alloys, etc.) and are subject to the QS/GMP. Manufacturers of human blood and blood components are not subject to this part, but are subject to 21 CFR 606.

The QS/GMP includes regulations regarding Purchasing Controls, 21 CFR 820.50. Receiving, In-process and Finished Device Acceptance, 21 CFR 820.80, and Traceability, 21 CFR 820.65, that require finished device manufacturers exercise more control over the components they use in their devices. The preamble of the QS/GMP states: “Since FDA is not regulating component suppliers, FDA believes that the explicit addition to the CGMP requirements of the purchasing controls...is necessary to provide the additional assurance that only acceptable components are used.” And “...inspections and tests, and other verification tools, are also an important part of ensuring that components and finished devices conform to approved specifications.” It further states, “...traceability of components must be maintained so potential and actual problem components can be traced back to the supplier.”
The medical device QS/GMP is an umbrella GMP that specifies general objectives rather than methods. It is left to the manufacturer to develop the best methods to meet these objectives. You must use good judgment in determining compliance with the QS/GMP, keeping in mind that it is an umbrella GMP and all requirements may not apply or be necessary. The purpose of the QS/GMP is to assure conformance to specifications and to ensure that all requirements that will contribute to assuring the finished device meets specifications are implemented. You should not insist that a manufacturer meet non-applicable requirements. Refer to IOM Exhibit 5-13 for types of establishments that are required to comply with the QS/GMP.

5.6.2.1 - Pre-Inspectional Activities

Prior to the start of any medical device inspection, the factory jacket or establishment history of the establishment should be reviewed. You should review the previous inspectional findings and subsequent correspondence between the establishment and FDA; any MDR or consumer complaints where it was determined follow-up would occur at the next inspection; and any notifications of recalls since the last inspection.

MDR data most useful in preparing for an inspection includes specific MDRs for the manufacturer (i.e., query by establishment's short name) for the time frame since the last inspection, or MDRs for the generic devices manufactured by that establishment (i.e., query by product code) for some reasonable time frame. This data assists you in determining potential problem areas in the manufacture or design of the device, or lot or batch specific issues. MDR information can be accessed through the electronic CDRH Information Retrieval System (eCIRS) and through Total Product Lifecycle Reports (TPLC).

The establishment's reported registration and listing data should be verified during any GMP inspection to assure there have been no changes and the registration and listing data was accurately reported. Changes or inaccuracies should be immediately reported to the district medical device registration and listing monitor. See also Field Management Directive (FMD) 92.

510(k) and PMA data assists you in determining what devices the establishment is manufacturing and whether any new devices have been designed or changed since the last inspection. This data is useful in focusing the inspection on new or changed devices as well as devices that are higher risk devices, i.e., Class II or III versus Class I. This information can be accessed through IMAGE 2000 plus and TPLC.

Since information about medical device firms is distributed listing, premarket, adverse event (MDR), and CDRH complaint details as well as high level recall and inspection information. ORA investigators can run these reports in order to prepare for inspections.

It is necessary to have access to both Business Objects and ORADSS in order to run EHR2 and TPLC reports. If you require access, send a request to the Employee Resource & Information Center (ERIC). You need to specify in your request to ERIC that you need access to the TPLC reports.

**Accessing EHR2 reports**

Go to http://inside.fda.gov:9003/it/Applications/ORAAplications/default.htm

Under ORADSS click go

Click the Documents Tab; then click Folders

Navigate (by hitting the plus sign) into Public Folders and then the Domestic Reports subfolder

Double click on the Establishment History Report subfolder

Select EHR02_Firm Info and double click

*Note: For initial inspections, there will not be any inspection history in the EHR2 report

**Accessing TPLC reports**

Go to http://bi.fda.gov

In the upper left corner, you will see tabs labeled “Home” and “Documents”. Click on “Documents”. At the bottom left, you will see tabs labeled “My Documents”, “Folders”, “Categories” and “Search”. Click on “Folders”. You may see a folder that is labeled “Public Folders”. Click on the folder. Navigate (by hitting the plus sign) into the DRLM Folder and then the Manufacturer and Product Details subfolder. The TPLC reports will appear to the right. Alternatively, if there is no plus sign next to your DRLM folder, you may see a separate TPLC folder further down on the list of folders. Click on TPLC. Select a report by double clicking the report name

Beneficial reports include:

- TPLC Manufacturer Name – displays information on a firm based on the Manufacturer name entered. The manufacturer name entered must be in upper case as the reports are case sensitive. Since company names vary with the inclusion of commas, abbreviations (INC vs. INC. vs. INCORPORATED), or division names, it is best to first use the shortest name possible with a wild card character (%). For example, to search for “XYZ SURGICAL CO (PVT) LTD”, it would be best to first use “XYZ%”. This will return all manufacturers that have a name beginning with that phrase. Given that some firms might have similar names, the report might return several companies with a name beginning with “XYZ”. The best option then is to look at those companies returned in
the report and modify the name used in the search.

- TPLC Product Code Reviewer - displays information on one or more specific product codes entered. Ensure that the product codes entered are in upper case as the reports are case sensitive.

If you have general questions or need help accessing this information, please contact your CDRH Field Inspection and Support Branch regional representative or MPTPOB at ORAHQDeviceInspectionPOC@fda.hhs.gov.

IOM 5.2 should be followed in regards to pre-announcement of medical device inspections.

5.6.2.2 - Quality Audit

The inspectional approach for identifying inadequate auditing of a quality assurance program is limited by the agency’s policy, which prohibits access to audit results. The policy is stated in CPG section 130.300 (7151.02). Under the QS/GMP regulation (21 CFR 820.180 (c)) this prohibition extends to evaluations or audits of suppliers, 21 CFR 820.50(a), and Management Reviews conducted per 21 CFR 820.20. Evidence of inadequate auditing may be discovered without gaining access to the written audit reports. See the Guide to Inspections of Medical Device Manufacturers or Guide to Inspections of Quality Systems for inspectional guidance.

The preamble to the QS/GMP specifically states, “FDA will review the corrective and preventive action procedures and activities performed in conformance with those procedures without reviewing the internal audit reports. FDA wants to make it clear that corrective and preventive actions, to include the documentation of these activities, which result from internal audits and management reviews are not covered under the exemption at 820.180(c).” Therefore, these corrective and preventive actions and documentation are not exempted from inspectional scrutiny.

The QS/GMP regulation (21 CFR 820.180(c)) requires a manufacturer to certify in writing that audits and reaudits have been conducted whenever requested to do so by an investigator. Investigators through their supervisors should consult with CDRH (HFZ-306) prior to requesting such certification.

5.6.2.3 - Records

FDA has distinct authority under section 704(e) of the FD&C Act [21 U.S.C. 374 (e)] to inspect and copy records required under section 519 or 520(g) of the FD&C Act [21 U.S.C. 360i or 360j (g)]. Investigators should only collect copies of documents as necessary to support observations or to satisfy assignments. Manufacturers who have petitioned for and obtained exemption from the QS/GMP are not exempted from FDA authority to review and copy complaints and records associated with investigation of device failures and complaints.

You may advise manufacturers they may mark as confidential those records they deem proprietary to aid FDA in determining which information may be disclosed under Freedom of Information.

Records must be maintained for as long as necessary to facilitate evaluation of any report of adverse performance, but not less than two years from the date the device is released for distribution. Records required by the Radiation Control for Health and Safety Act must be maintained for five years. It is permissible to retain records in electronic or photocopy form, providing the copies are true and accurate reproductions.

5.6.2.4 - Complaint Files

Complaints are written or oral expressions of dissatisfaction with finished device identity, quality, durability, reliability, safety, effectiveness or performance. Routine requests for service would not normally be considered complaints. However, service requests should be reviewed to detect complaints, and as part of any trend analysis system, and to comply with 820.20(a)(3).

FDA has the authority to require a device firm to open its complaint files, and review and copy documents from the file.

Provisions in the FD&C Act pertaining to FDA review of records are:

1. For restricted devices the FD&C Act in Section 704(a)(1)(B) [21 U.S.C. 374 (a)(1)(B)] extends inspection authority to records, files, papers, processes, controls and facilities bearing on restricted medical devices. See FD&C Act Sec. 704 [21 U.S.C. 374] for a full explanation and for a list of the items, e.g., financial data, which are exempt from disclosure to FDA.

2. For all devices, including restricted devices, refer to Section 704(e) of the FD&C Act [21 U.S.C. 374 (e)], which provides for access to, copying and verification of certain records.

3. Section 519 of the FD&C Act [21 U.S.C. 360i] requires manufacturers, importers, or distributors of devices intended for human use to maintain such records, and provide information as the Secretary may by Regulation reasonably require.

4. Section 520(g) of the FD&C Act [21 U.S.C. 360j (g)] covers the establishment of exemptions for devices for investigational use and the records which must be maintained and open for inspection.

QS/GMP requirements for complaint files are found in 21 CFR 820.198. GMP requirements for complaint files first became effective on December 18, 1978. The Quality System Regulation, which went into effect on June 1,
1997, added to and modified the requirements for complaint handling. The regulation contains a provision that records maintained in compliance with the QS/GMP must be available for review and copying by FDA (21 CFR 820.180). Complaint files are QS/GMP required records; therefore, the manufacturer must make all complaints received on or after December 18, 1978 and the records of their investigation available for FDA review and copying. EIRs should contain enough information to allow cross-referencing between complaints and MDRs.

21 CFR Part 803 requires medical device manufacturers to report deaths, serious illnesses, and serious injuries to FDA for which a device has or may have caused or contributed, and manufacturers must also report certain device malfunctions. The MDR reportable events must be maintained in a separate portion of the complaint files or otherwise clearly identified. These complaints must be investigated to determine whether the device failed to meet specifications; whether the device was being used for treatment or diagnosis; and the relationship, if any, of the device to the reported incident or adverse event.

When a firm determines complaint handling will be conducted at a place other than the manufacturing site, copies of the record of investigation of complaints must be reasonably accessible at the actual manufacturing site.

### 5.6.3 - STERILE DEVICES

Inspections of sterile device manufacturers are conducted per Compliance Program Guidance Manual 7382.845, as a production process under the Production and Process Control Subsystem. See the Guide to Inspections of Quality Systems for further guidance.

### 5.6.4 - LABELING

Specific labeling requirements for in vitro diagnostics (IVDs) are contained in 21 CFR 809.10.

Part 809.10(a) contains explicit labeling requirements for the individual IVD container, and for the outer package labeling and/or kit labeling. Part 809.10(b) contains special labeling requirements for the product insert, which must be included with all IVD products. These two sections also contain the requirements for: lot numbers, allowing traceability to components (for reagents) or subassemblies (for IVD instruments); stability studies for all forms of the product; an expiration date, or other indication to assure the product meets appropriate standards; and, the requirements for establishing accuracy, precision, specificity and sensitivity (as applicable).

Part 809.10(c) lists the labeling statements required for IVDs which are being sold for investigational and research use. Determine whether the firm is limiting the sale of IVDs, labeled as such, to investigators or researchers. Document any questionable products, and submit to CDRH for review.

Warning and caution statements recommended for certain devices, along with certain restrictions for use, are described in 21 CFR 801. This same section also contains the general labeling regulations, which apply to all medical devices.

### 5.6.5 - GOVERNMENT-WIDE QUALITY ASSURANCE PROGRAM (GWQAP)

Inspections under the GWQAP are conducted upon request by Office of Enforcement and Import Operations (OEIO), Division of Compliance Systems (DCS). Each assignment is specific and may involve more than a single compliance program. Specific questions arising during or as a result of these inspections should be directed to OEIO/DCS.

### 5.6.6 - CONTRACT FACILITIES

Device manufacturers may employ the services of outside laboratories, sterilization facilities, or other manufacturers (i.e., injection molders, packagers, etc.). The finished device manufacturer is responsible for assuring these contractors comply with the QS/GMP and that the product or service provided is adequate. These contractors are subject to FDA inspection and some are subject to the QS/GMP regulation. This “...includes but is not limited to those who perform the functions of contract sterilization, installation, relabeling, remanufacturing, repacking, or specification development, and initial distributors of foreign entities performing these functions,” per 21 CFR 820.3(o). Whether under contract or not if a firm manufactures a finished device by the definition found in 21 CFR 820.3(l) “Finished device” means any device or accessory to any device that is suitable for use or capable of functioning, whether or not it is packaged, labeled, or sterilized they are subject to QS/GMP. NOTE: if the product manufactured by the contractor also meets the definition of a component and a finished device, the contractor is subject to the QS/GMP regulation.

Determine how a manufacturer evaluates and selects potential contractors for their ability to meet the manufacturer’s requirements, as required by 820.50, Purchasing Controls. Conducting audits can be an effective method for assessment. However, not all contractors allow audits. Audits may not be feasible in some instances. In other instances the activity the contractor is conducting may not have a significant impact on the device safety or function; therefore, expending the resources necessary to audit the contractor may not be warranted.

Evaluations may be accomplished by other means such as requesting that the potential contractor fill out a questionnaire about their quality system, asking other customers of the contractor about their experiences with the firm, or basing assessments on past performance. Evaluations must be documented. The extent to which a manufacturer has evaluated a contractor, as well as the results of the evaluation, should govern the degree of
oversight exercised over products and services supplied by the contractor.

5.6.7 - SMALL MANUFACTURERS

When inspecting one-person or very small manufacturers for compliance with the QS/GMP master record and written procedure requirements, the investigator should realize that detailed written assembly, process, and other instructional procedures required for larger firms may not be needed. In a small firm, division of work is at a minimum, with one person often assembling and testing the finished device. In many cases, blueprints or engineering drawings could be adequate procedures. The QS regulation requires that certain activities be defined, documented and implemented. The regulation does not require separate procedures for each requirement and often several requirements can be met with a single procedure. The complexity of the procedures should be proportional to the complexity of the manufacturer's quality system, the complexity of the organizational structure and the complexity/risk of the finished device being produced. In assessing the need for detailed or lengthy written procedures, the investigator should make judgments based on training and experience of the individuals doing the work and the complexity of the manufacturing process. However, this does not mean small manufacturers have any less responsibility for complying with the QS regulation or assuring safe and effective devices are produced.

5.6.8 - BANNED DEVICES

Section 516 of the FD&C Act [21 U.S.C. 360f] provides a device for human use may be banned by regulation (21 CFR 895) if it presents substantial deception or an unreasonable and substantial risk of illness or injury. Investigators should become familiar with this regulation. When you determine, during an inspection or investigation, that banned devices are being distributed, the distribution, manufacture, etc., should be documented as for any other violative product.

5.6.9 – REPORTS OF CORRECTIONS AND REMOVALS

Manufacturers, importers, and distributors of medical devices are to promptly report to the FDA any corrections or removals of a device undertaken to reduce a risk to health posed by the device or to remedy a violation of the FD&C Act caused by the device which may present a risk to health as provisioned by the Safe Medical Devices Act of 1990 and 21 CFR Part 806. Refer to IOM Ch. 7 – RECALL ACTIVITIES 7.2.3 MEDICAL DEVICE RECALLS for more information.

5.6.10 – TRACKED MEDICAL DEVICES

A "tracked medical device" is a device regulated by CDRH and for which the firm has received "tracking orders". CDRH has a dedicated mailbox to manage inquiries about tracked devices and the related regulation at TrackedDevicesMailbox@FDA.HHS.GOV.

5.6.11 - DEVICE INSPECTION REPORTS

See IOM 1.1, English language requirement. You should write your EIR following the guidance in IOM 5.10.4, 5.10.4.1, 5.10.4.2, 5.10.4.3. Section headings can be added to address the needs of other Compliance Program Guidance Manuals such as 7383.001 for pre-market and post-market PMA inspections. Include in your report the systems, processes, products, and product classification covered during the current inspection.

SUBCHAPTER 5.7 - BIOLOGICS

5.7.1 - DEFINITION

A "biological product" means a virus, therapeutic serum, toxin, antitoxin, vaccine, blood, blood component or derivative, allergenic product, or analogous product, or arsphenamine or derivative of arsphenamine (or any other trivalent organic arsenic compound), applicable to the prevention, treatment, or cure of a disease or condition of human beings (Public Health Service Act Sec. 351(i)). Additional interpretation of the statutory language is found in 21 CFR 600.3. Biological products also meet the definition of either a drug or device under Sections 201(g) and (h) of the Federal Food, Drug, and Cosmetic Act (FD&C Act).

Veterinary biologicals are subject to the animal Virus, Serum, and Toxin Act which is enforced by USDA (21 U.S.C. 151-158).

5.7.2 - BIOLOGICS INSPECTIONS

FDA has developed a strategy known as "Team Biologics", a reinvention of the agency's approach to inspectional coverage of certain biological products. The periodic CGMP inspections and compliance operations of plasma fractionated products, allergenic products, vaccines, gene and cell therapy products, and biological in vitro diagnostic devices are led by investigators and compliance officers on Team Biologics. Team Biologics investigators and its compliance officers report to DMPTPO. Inspections of unlicensed CBER-regulated medical devices are not covered by the Team (e.g., blood establishment software) but are conducted by District investigators who may or may not be part of the Biologics Cadre. See IOM 2.2 for a discussion of statutory authority. CBER maintains the lead for pre-licensing and pre-approval inspections of biological products, while ORA customarily leads PMA/510(k) inspections.

5.7.2.1 - Authority

Biological products are regulated under the authority of Section 351 of the Public Health Service Act and under the Food, Drug, and Cosmetic Act, as drugs or devices, with the exception of certain human cells, tissues, and
cellular and tissue-based products (HCT/Ps) regulated solely under Section 361 of the Public Health Service Act (see 21 CFR 1271.10). Blood and blood products for transfusion are prescription drugs under the FD&C Act. Under the FD&C Act, source plasma and recovered plasma may have the legal identity of either a drug or device depending on its intended use. Section 351(a) of the PHS Act provides for licensure of biological products and inspection of the products covered is per 351(d). Most biological drugs are licensed. The investigational new drug application regulations (21 CFR 312) also apply to biological products subject to the licensing provisions of the PHS Act. However, investigations of blood grouping serum, reagent red blood cells, and anti-human globulin in-vitro diagnostic products may be exempted (21 CFR 312.2(b)).

### 5.7.2.1.1 - BLOOD AND SOURCE PLASMA INSPECTIONS

The investigators in the Biologics Cadre perform inspections of blood and plasma establishments. For blood bank and source plasma establishments, use the CGMPs for Blood and Blood Components (21 CFR 600) as well as the general requirements for biological products (21 CFR Part 600), the general biological product standards (21 CFR Part 610), and the additional standards for human blood and blood products (21 CFR Part 640.) The drug GMPs (21 CFR 200/211) also apply to biological drugs. In the event it is impossible to comply with both sets of regulations, the regulation specifically applicable to the product applies. This would generally be Parts 606 and 640 of the regulations in the case of blood bank and source plasma establishments.

### 5.7.2.1.2 – HUMAN TISSUE INSPECTIONS

In February 1997, FDA proposed a new, comprehensive approach to the regulation of human cellular and tissue-based products (now called human cells, tissues, and cellular and tissue-based products or HCT/Ps). The agency announced its plans in two documents entitled, "Reinventing the Regulation of Human Tissue" and "A Proposed Approach to the Regulation of Cellular and Tissue-based Products" (62 FR 9721, March 4, 1997).

Since that time, the agency has published three final rules and one interim final rule to fully implement the proposed approach. On January 19, 2001, FDA finalized regulations to create a new, unified system for registering HCT/P establishments and for listing their HCT/Ps (registration final rule, 66 FR 5447). Part of the definition of "human cells, tissues, or cellular or tissue-based products" became effective on January 21, 2004. On January 27, 2004 (69 FR 3823), we issued an interim final rule to except human dura mater and human heart valve allografts from the scope of that definition until all of the tissue rules became final. On May 25, 2004, FDA finalized regulations requiring most cell and tissue donors to be tested and screened for relevant communicable diseases (donor-eligibility final rule, 69 FR 29786). On November 21, 2004, FDA finalized regulations requiring HCT/P establishments to follow current good tissue practice (CGTP), which governs the methods used in, and the facilities and controls used for, the manufacture of HCT/Ps; recordkeeping; and the establishment of a quality program. The new CGTP regulations also contain certain labeling and reporting requirements, as well as inspection and enforcement provisions (GTP final rule, 69 FR 68612). The donor eligibility and CGTP rules became effective May 25, 2005.

Part 1271 contains six subparts:

1. Subpart A of part 1271 – general provisions
2. Subpart B of part 1271 - registration
3. Subpart C of part 1271 - screening and testing of donors to determine eligibility
4. Subpart D of part 1271 - provisions on CGTP
5. Subpart E of part 1271 - certain labeling and reporting requirements

The subparts apply as follows:

Subparts A through D apply to all HCT/Ps, i.e., to those HCT/Ps described in Sec. 1271.10 and regulated solely under section 361 of the PHS Act, and to those regulated as drugs, devices, and/or biological products. Subparts E and F, which pertain to labeling, reporting, inspection, and enforcement, apply only to those HCT/Ps described in Sec. 1271.10 and regulated solely under section 361 of the PHS Act. However, with the exception of two provisions (Sec. 1271.150(c) and 1271.155) subparts D and E are not being implemented for reproductive HCT/Ps described in 21 CFR 1271.10 and regulated solely under section 361 of the PHS Act.

HCT/Ps subject to the provisions of 21 CFR Part 1271 include, but are not limited to, bone, ligaments, skin, dura mater, heart valve, cornea, hematopoietic stem/progenitor cells derived from peripheral and cord blood, manipulated autologous chondrocytes, epithelial cells on a synthetic matrix, and semen or other reproductive tissue.

For HCT/P inspections, use the CP 7341.002, “Inspections of Human Cells, Tissues, and Cellular and Tissue-Based Products.”

### 5.7.2.2 – Donor Confidentiality

Blood bank, source plasma, and human tissue establishments are sensitive to maintaining confidentiality of donor names. The mere reluctance to provide records is not a refusal. However, FDA has the authority under Section 704 of the FD&C Act to make inspections and 21 CFR 600.22(g) and 1271.400(d) provides for copying records during an establishment inspection. For prescription drugs, section 704 of the FD&C Act...
specifically identifies records, files, papers, processes, controls, and facilities as being subject to inspection.

If you encounter problems accessing records, explain FDA's authority to copy these records. IOM 5.2.5 should be followed if a refusal is encountered. When donor names or other identifiers are necessary, they may be copied, but the information must be protected from inappropriate release. See IOM 5.3.8.6.

5.7.2.3 – Inspecti onal Objectives

The inspectional objective for biological products is to assure the products are safe, effective, and contain the quality and purity they purport to possess, and are properly labeled. The inspectional objective for HCT/Ps is to assure that HCT/Ps are recovered, processed, stored, labeled, packaged and distributed, and the donors are screened and tested, in a way that prevents the introduction, transmission, or spread of communicable diseases. Facilities will be inspected for conformance with:

1. Provisions of the PHS Act and FD&C Act,
3. HCT/P regulations in 21 CFR 1270 and 1271.
4. FDA Policies, which include guidance to the industry, and the Compliance Policy Guides Chapter 2.

5.7.2.4 - Preparation

Review the district files of the facility to be inspected and familiarize yourself with its operation and compliance history. Review:

1. Appropriate Compliance Programs and related Compliance Policy Guides (CPG), Chapter 2.
   NOTE: Federal Cooperative Agreements Manual; MOU with the Department of Defense, and MOU with the Centers for Medicare and Medicaid Services (CMS) on transfusion services;
2. Correspondence from the firm depicting any changes since the last inspection;
3. Firm's registration and product listing information;
5. Biological Product Deviation Reports, Adverse Reaction Reports, complaints, and recalls;

Through guidance documents, CBER sets forth its inspection policy and regulatory approach. A list of these documents is attached to the current Compliance Program Guidance Manuals (CP) available on the CBER internet site at (CBER CP Website).

The OSHA regulation 29 CFR 1910.1030 dated December 6, 1991, was intended to protect health care workers from blood borne pathogens, including those involved in the collection and processing of blood products. The regulation defines expectations for the use of gloves, hand washing facilities, decontamination of work areas, waste containers, labeling and training of employees and exemptions for volunteer blood donor centers. FDA Investigators should adhere to these safety guidelines during inspections or related activities in establishments that process biologically hazardous materials.

Become familiar with the OSHA regulations and their applicability to 21 CFR 606.40(d)(1) and (2), which require the safe and sanitary disposal for trash, items used in the collection and processing of blood and for blood products not suitable for use. Consult your district biologics monitor for copies of the above references. Additional copies may be obtained from OO, OMPTO, Division of Medical Products and Tobacco Program Operations or see CBER's web site.

5.7.2.5 - Inspectional Approach

Use the Compliance Program Guidance Manuals (CP) and Guides to Inspection of Source Plasma Centers and Infectious Disease Marker Testing Facilities for inspectional instructions. The EIR must clearly identify the areas covered. The report should include a summary of the inspection, the FDA 482, the FDA 483, if issued, and the required FACTS EI Record.

Particular attention should be given to biological products deviation reports indicative of problematic areas or processes, adverse reactions, transfusion associated AIDS (TAA), transfusion or donation associated fatalities and hepatitis and HIV lookback procedures. For additional information regarding TAA, see CP 7342.001. The follow-up investigations to such reports should also be covered.

Complaints, in particular those involving criminal activity, must be promptly investigated and coordinated with other agency components as needed.

For blood banks and source plasma establishments, refer to CP 7342.001 and 7342.002 for a discussion of the systems approach to inspection. The CP incorporates a systems-based approach to conducting an inspection and identifies five (5) systems in a blood bank and source plasma establishment operation for inspection. Each system may not be in a particular establishment operation; therefore, the inspection should focus on the systems present. The CP directs an in-depth audit of the critical areas in each system. A multi-layered system of safeguards has been built into the blood collection, manufacturing and distribution system to assure a safe blood supply.

For HCT/P establishments, refer to CP 7341.002.

For Biological Drug Products, refer to CP 7345.848.

For Licensed Viral Marker Test Kits, refer to CP 7342.008.
If Investigators encounter products not specifically referenced in the regulations, they should contact CBER/OCBQ/Division of Inspections and Surveillance for guidance.

**5.7.2.6 - Regulations, Guidelines, Recommendations**

Guidance documents for industry are made available to the public in accordance with good guidance practice regulations at 21 CFR 10.115. The contents of most of these documents are incorporated into the establishment's SOPs and/or license applications or supplements. In addition, DMPTPO has issued Source Plasma Establishment and Infectious Disease Marker Testing Facility Inspectional Guides to be used by investigators during inspections.

Deviations from guidance documents must not be referenced on a FDA 483. However, since these documents are often related to specific GMP requirements, in most cases deviations can be referenced back to the GMP. If a deviation is observed during an inspection and the investigator relates it to the regulations or law, then the item may be reported on the FDA 483. During the discussion with management, the relationship of the deviation to the regulation or law, or accepted standard of industry, should be clearly explained.

If an establishment indicates it is not aware of any of these documents, provide the telephone number of CBER's Office of Communication, Outreach and Development, Division of Training, and Manufacturers Assistance, 301-827-2000.

If a firm claims approval for an alternative procedure, verify by reviewing the firm's written approval letter. Approved alternative procedures may be verified by contacting CBER/Division of Blood Applications or the appropriate CBER product office.

**5.7.2.7 - Technical Assistance**

Several FDA regions and districts have biologics specialists who are available for technical assistance and consultation. Do not hesitate to avail yourself of their services.

The services of expert investigators in ORA/OO/OMPTO/DMPTPO, 301-796-0358, are available for telephone or on-site consultation and assistance in problem areas.

CBER/OCBQ, Division of Inspections and Surveillance (HFM-650), 301-827-6220, can provide technical assistance, and can coordinate assistance with other CBER offices.

**5.7.2.8 - CBER Bioresearch Monitoring**

Bioresearch monitoring (BIMO) assignments for biological products will generally be issued by the Center for Biologics Evaluation and Research (CBER) (see IOM 5.5.6).

**5.7.3 - REGISTRATION, LISTING AND LICENSING.**

**5.7.3.1 - Registration and Listing**

See IOM 2.9.3.1

**5.7.3.1.1 – TRANSFUSION SERVICES**

Most transfusion services are exempt from registration under 21 CFR 607. This includes facilities that are certified under the Clinical Laboratory Improvement Amendments of 1988 (42 U.S.C. 263a) and 42 CFR Part 493 to perform the FDA-required tests on blood or has met equivalent requirements as determined by the Centers for Medicare and Medicaid Services, and are engaged in the compatibility testing and transfusion of blood and blood components, but which neither routinely collect nor process blood and blood components. Such facilities include establishments:

1. Collecting, processing and shipping blood and blood components under documented emergency situations,
2. Performing therapeutic phlebotomy and therapeutic plasma exchange after which the product is discarded,
3. Preparing recovered human plasma and red blood cells,
4. Pooling products/platelets for in-house transfusion,
5. Thawing frozen plasma or cryoprecipitate for transfusion.

**5.7.3.1.2 - HCT/PS**

Establishments manufacturing HCT/Ps (human cells, tissues, or cellular or tissue-based products) as defined in 21 CFR 1271.3(d) must register and list using form FDA 3356. Examples of HCT/Ps include, but are not limited to, bone, ligament, skin, cornea, hematopoietic stem cells derived from peripheral and cord blood, manipulated autologous chondrocytes, and semen or other reproductive tissue. Establishments manufacturing HCT/Ps regulated as medical devices, drugs or biological drugs must also register and list with the FDA pursuant to 21 CFR 1271 using form FDA 3356.

**5.7.3.1.3 - LABORATORIES**

Laboratories performing infectious disease testing of donors of blood or blood components or HCT/P are an FDA obligation and required to register. Clinical laboratories were previously exempted from registration by 21 CFR 607.65(g), but FDA revoked this regulation. Your inspections should focus on activities relevant to blood product and HCT/P testing operations.

**5.7.3.1.4 - MILITARY BLOOD BANKS**
CHAPTER 5 INVESTIGATIONS OPERATIONS MANUAL 2016

Inspection of military blood banks is an ORA responsibility. These facilities are required to meet the same standards as other blood banks although military emergencies may require deviations from the standards. A separate license is held by each branch of the service; although each individual establishment may be licensed or unlicensed, all are required to register. Districts should notify the appropriate military liaisons 30 days before inspection of a military facility. For additional information on inspection of government establishments, see Compliance Program Guidance Manual 7342.001, the Federal Cooperative Agreements Manual, and the MOU with Department of Defense Regarding Licensure of Military Blood Banks.

Field Management Directive 92, Agency Establishment Registration and Control Procedures, details the registration process within the agency.

Ensure the firm’s current registration forms reflect actual operations.

5.7.3.2 – MOUs

Under the 1983 Memorandum of Understanding (MOU) between the FDA and the Centers for Medicare and Medicaid Services (CMS, formerly Health Care Financing Administration - HCFA), CMS agreed to survey these facilities that engage in minimal manufacturing in order to minimize duplication of effort and reduce the burden on the affected facilities while continuing to protect transfusion recipients. However, no transfer of statutory functions or authority is made under the MOU and the FDA retains legal authority to inspect these unregistered transfusion services whenever warranted. When appropriate, Districts should conduct inspections jointly with the CMS regional liaison. If you determine during a routine inspection an establishment is a CMS obligation under the MOU, you should terminate the inspection and report as such. See Federal Cooperative Agreements Manual - FDA/HCFA Memorandum of Understanding.

5.7.3.3 - Biologic License

See IOM 2.9.3.2. A biologics license application (BLA) shall be approved only after inspection of the establishment(s) listed in the application and upon a determination that the establishment complies with the standards established in the BLA and the requirements prescribed in applicable regulations (21 CFR 601.20(d)). CBER is responsible for conducting all pre-license (PLI) and pre-approval (PAI) inspections of CBER-regulated products. These inspections are part of the review of a BLA or BLA supplement. CBER identifies the scope of the inspection and invites ORA to participate in the inspections. Copies of CBER’s PLI and PAI inspection reports are forwarded to the districts and should be part of the firm’s file.

5.7.3.4 - Approval of Biological Devices

There must be a pre-approval inspection (PAI) of the establishment for compliance with the QS/GMP regulation and the firm’s PMA. For licensed devices, CBER conducts the pre-license inspection (PLI). Devices used in the collection and testing of blood for transfusion are approved/cleared through the PMA/510(k) authorities. ORA Investigators customarily inspect the CBER regulated devices, which are subject to PMA/510(k) applications.

5.7.4 - RESPONSIBLE INDIVIDUALS

In licensed establishments, the applicant or license holder may designate an authorized official(s) to represent the applicant to the FDA in matters of compliance. The FDA 482 and any 483 should be issued to the most responsible person on the premises at the time of inspection. An exact copy of the FDA 483 should also be forwarded to the top official of the firm if that person did not receive the FDA 483. The designation as authorized official does not necessarily mean that individual is the most responsible for any non-compliance of the firm. In licensed or unlicensed facilities, establish and document all individuals responsible for violations and their reporting structure in the organization.

5.7.5 - TESTING LABORATORIES

Blood bank, source plasma, and HCT/P establishments may use outside testing laboratories to perform required testing.

Laboratories conducting testing for licensed blood banks are usually licensed. CBER may approve the use of a non-licensed laboratory to do required testing, provided the lab is capable of performing the tests and the lab registers with CBER prior to CBER approving the licensing arrangement.

Laboratories performing required testing for source plasma manufacturers must either be:
1. Licensed or
2. Certified to perform such testing on human specimens under the Clinical Laboratory Improvement Amendments of 1988 (42 U.S.C. 263a) and 42 CFR part 493, or has met equivalent requirements as determined by CMS.

Laboratories performing required testing for HCT/Ps must:
1. Test using approved FDA-licensed, approved or cleared donor screening tests according to the manufacturers instructions, and
2. Be either certified to perform such testing on human specimens under the Clinical Laboratory Improvement Amendments of 1988 (42 U.S.C. 263a) and 42 CFR part 493, or has met equivalent requirements as determined by CMS.

Instructions for inspecting testing laboratories are included in the appropriate Compliance Program Guidance Manuals. Coordinate the inspection of non-registered laboratories with CMS regional office contacts. If a testing
laboratory is located outside of the district, request an inspection by the appropriate district office, where appropriate.

5.7.6 - BROKERS

Blood establishments may use brokers to locate buyers for products such as recovered plasma or expired red blood cells. These articles are used for further manufacture into products such as clinical chemistry controls and in-vitro diagnostic products not subject to licensure. Fractionators also use brokers to locate suppliers of plasma under the short supply provisions (21 CFR 601.22). During inspections, determine if the facility is selling products to any brokers. If brokers are used, determine if the brokered products are shipped to a facility operated by the broker or directly to the consignee.

Brokers who take physical possession of blood products and engage in activities considered manufacturing or labeling are required to register and are included in the OEI for routine inspection under the blood bank compliance program. Brokers who only arrange sales of or store blood and blood components, but do not engage in manufacturing activities are not required to register.

SUBCHAPTER 5.8 - TOBACCO PRODUCTS

5.8.1 - DEFINITIONS

The term "tobacco product" is defined in FD&C Act Section 201(rr) and means any product made or derived from tobacco that is intended for human consumption, including any component part, or accessory of a tobacco product (except for raw materials other than tobacco used in manufacturing a component, part, or accessory of a tobacco product.) The term "tobacco product" does not mean an article that is a drug under section 201(g)(1) of the FD&C Act, a device under section 201(h) of the FD&C Act, or a combination product described in section 503(g) of the FD&C Act.

The definition of certain tobacco products can be found in the FD&C Act under section 900.

5.8.2 – TOBACCO INSPECTIONS

See IOM 2.2 for discussion of statutory authority

For the first few years, inspections involving tobacco product(s) at manufacturing facilities should be made pursuant to an assignment until a Compliance Program is developed. CTP’s office of Compliance and Enforcement and ORA’s Division of Medical Products and Tobacco Program Operations are available to work with the field during inspections.

5.8.3 - RETAIL COMPLIANCE CHECK INSPECTION CONTRACTS

FDA issues contracts to assist with compliance check inspections of retail establishments to help enforce the Youth Access and Advertising Regulations that took effect on June 22, 2010. FDA has a goal of establishing a contract, where feasible, with every U.S. State and Territory, but some States and Territories, for a variety of reasons, have been unable to do so. Therefore, FDA has awarded contracts to third-party entities that are able to hire commissionable inspectors to do compliance check inspections of tobacco retailers in those states and territories where FDA has not been able to contract with a state agency. FDA has further expanded this program by awarding retail inspection contracts to Tribes to conduct retail inspections within their jurisdictions.

5.8.4 - GUIDANCE, COMPLIANCE & REGULATORY INFORMATION

The Center for Tobacco Products website contains resources for legal, regulatory, and policy issues related to tobacco products and information for small business assistance (SmallBiz.Tobacco@fda.hhs.gov).

SUBCHAPTER 5.9 - VETERINARY MEDICINE

5.9.1 - CVM WEBSITE

The Center for Veterinary Medicine website contains; a listing of current and planned Guidance Documents; and on-line access to the Animal Drug@fda database listing new animal drug approvals. There is a “search” feature allowing you to search for documents containing various words or phrases. The website also contains organizational information for the Center and an explanation of the various laws and regulations which the Center enforces. Information on the website can provide guidance for inspectional efforts related to CVM obligations.

5.9.2 - VETERINARY DRUG ACTIVITIES

CVM is responsible for inspections of therapeutic and production drugs, and Active Pharmaceutical Ingredients (APIs). Therapeutic drugs are used in the diagnosis, cure, mitigation, treatment or prevention of disease. Production drugs are used for economic enhancement of animal productivity. Examples include: growth promotion, feed efficiency and increased milk production.

Preapproval inspections are conducted pursuant to pending NADA or ANADA applications.

Post approval inspections of veterinary drugs are conducted to determine compliance with the Current Good Manufacturing Practices (CGMPs) for Finished Pharmaceuticals under 21 CFR Part 211. These cGMPs apply to both human and veterinary drugs. Information on
veterinary drugs approved can be found in the "Green Book" database accessed through CVM's website.

APIs are active pharmaceutical ingredients. Many of the APIs used to manufacture dosage form drugs are imported from foreign countries. The intended source for an API must be indicated in NADA/ANADA submissions for new animal drug approvals. Any change in a source for an API would require a supplement to the application.

Extra label drug use refers to the regulations in 21 CFR Part 530 codified as a result of the Animal Medicinal Drug Use Clarification Act (AMDUCA) of 1994. These regulations set forth the requirements that veterinarians must meet to prescribe extra label uses of FDA approved animal and human drugs. The regulations describe what is a valid veterinary-client-patient relationship as well as what is considered illegal extra label use. 21 CFR Part 530 addresses issues regarding extra label use in non-food as well as food producing animals. 21 CFR 530.41 contains a list of drugs that cannot be used in an extra label manner in food-producing animals. During an inspection or investigation if you encounter any situations on suspected illegal extra label use of any FDA approved animal or human drugs or those prohibited for extra label use in food animals, you should contact CVM's Division of Compliance (HFV-230) (240-276-9200).

21 CFR Part 530 also addresses compounding of products from approved animal or human drugs by a pharmacist or veterinarian. The regulations clearly state compounding is not permitted from bulk drugs. This would include APIs. CVM has an existing CPG on Compounding of Drugs for Use in Animals (CPG 608.400). A copy can be found on CVM's website. The Division of Compliance (HFV-230) has issued assignments to conduct inspections of firms, including internet pharmacies, who may be engaged in the practice of manufacturing under the guise of pharmacy compounding. You should contact the Division of Compliance (HFV-230) at 240-276-9200 to report instances of compounding or to seek guidance on inspectional issues, or regulatory and enforcement policies.

Extra label drug use of any FDA approved animal or human drugs. The regulations clearly state what is a valid veterinary-client-patient relationship as well as what is considered illegal extra label use. 21 CFR Part 530 addresses issues regarding extra label use in non-food as well as food producing animals. 21 CFR 530.41 contains a list of drugs that cannot be used in an extra label manner in food-producing animals. During an inspection or investigation if you encounter any situations on suspected illegal extra label use of any FDA approved animal or human drugs or those prohibited for extra label use in food animals, you should contact CVM's Division of Compliance (HFV-230) (240-276-9200).

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5.9.3 - MEDICATED FEEDS AND TYPE A ARTICLES

Animal feed is defined under section 201(w) of the FD&C Act [21 U.S.C. 321 (w)]. CVM is responsible for control of medicated and non-medicated animal feeds, Type A medicated articles and pet foods.

The regulations for animal food labeling are in 21 CFR Part 501. The regulations for medicated feed mill licensure are in 21 CFR Part 515. The cGMPs for Medicated Feeds are in 21 CFR Part 225. The cGMPs for Type A Articles are in 21 CFR Part 226.

Inspections are routinely conducted of medicated feed mills and manufacturers of Type A Medicated Articles.

If you have questions related to cGMPs and enforcement policies and strategies concerning Medicated Feeds and Type A Articles you should contact the CVM/Division of Compliance (240-276-9200).

Guidance on pet food labeling requirements can be found on CVM's website.

5.9.4 - BSE ACTIVITIES

CVM is responsible for FDA's industry education and regulatory activities involving BSE and animal feed. BSE is "Bovine Spongiform Encephalopathy" and is often referred to as "mad cow disease." There are two BSE-related feed regulations: 21 CFR 589.2000, entitled "Animal Proteins Prohibited in Ruminant Feed" which was adopted in 1997, addresses the feeding of ruminant animals. A second rule, 21 CFR 589.2001, entitled "Cattle Materials Prohibited in Animal Food or Feed to Prevent the Transmission of Bovine Spongiform Encephalopathy" was adopted in 2009. 21 CFR 589.2001 prohibits the use of certain cattle-origin materials in the feed of all animals and is aimed primarily at rendering operations.

CVM has Guidance Documents in place dealing with BSE. The guidance documents address renderers, protein blenders, feed manufacturers, distributors and on farm feeders. The Compliance Program Guidance Manual and the inspection checklist are available on the CVM website, as are a variety of other BSE information, including a database containing a summary of the most recent inspection of each firm.

Questions on inspectional assignments and regulatory activities in the BSE area should be addressed to the CVM/Division of Compliance (HFV-230) at 240-276-9200.

5.9.5 - TISSUE RESIDUES

The presence of violative drug residues in food from slaughtered animals is a human health concern. Tissue residue inspections are performed in response to reports of violative drug residue levels found in tissue sampled at slaughter by the USDA/Food Safety Inspection Service (FSIS).

Tissue residues are commonly caused by medicating animals prior to marketing and failure to follow the drug's approved label directions. When a new animal drug is approved the approval is very specific in how the drug should be used, the dosage it should be given, route of administration, frequency of use and reason for use. A drug manufacturer conducts studies to determine withdrawal times and these times must be followed. Established tolerances for drug residues of new animal drugs in food can be found in 21 CFR Part 556.
Tissue residue investigations are unique in comparison to other fieldwork. Although your investigation may begin at the USDA slaughter establishment or person named on the USDA/FSIS “Violation Notification Letter,” you may inspect and/or visit more sites as part of your overall investigation. You may have to visit an auction barn, dealer, trucker, veterinarian, drug supplier, slaughter facility (USDA firm management or State personnel), etc. One or more of these establishments may be responsible for the tissue residue. Thus, each establishment’s activities may warrant a recommendation for regulatory action such as Warning Letter, Injunction, etc. when involvement with residue violations is documented.

Upon receipt of a FACTS assignment from CVM to conduct a tissue residue follow-up investigation, the district may also create additional operations, linked to the original CVM assignment, which will include all operations required to complete the CVM assignment. This could include multiple inspections, sample collections and/or investigations. You may not be aware of all the establishments you will visit prior to beginning your investigation. Appropriate operations should be added to or deleted from the district assignment.

Each site visit is unique and each produces its own set of unique documents and evidence requiring individual reporting by establishment. You should use good judgment during case development to assure you document your investigation thoroughly. Explain the chain of events and evidence, from the initial tissue residue report, and how other establishments were involved. Collect samples (usually DOC samples) as appropriate. Consultation with your supervisor and/or compliance branch during these operations is essential to assure all evidence necessary to develop a quality case is obtained and submitted in an appropriate format.

Following completion of all operations, you should prepare a Memo of Investigation referencing the FACTS assignments for your supervisor’s endorsement to the district Compliance Branch, with a copy to the originating CVM office. This Memo will summarize each site visit (EI or Investigation), sample(s) collected and relevance to the overall CVM assignment. A copy of the memo will be routed to each appropriate factory file.

The individual operations will then stand alone and/or may be used together to build one or multiple cases.

For example, a site visit to a slaughter facility may obtain information on the animal from the USDA inspection personnel on site; and obtain verification from management the establishment ships in interstate commerce. Information obtained at the slaughter facility or other establishments may be documented in an affidavit from each individual providing salient information. A site visit to a veterinarian may be important to establish whether the drugs which caused the tissue residue(s) were prescribed and, if so, how they were prescribed.

When there is reason to believe off-label use or other activities have occurred which may warrant a recommendation for regulatory action, an establishment inspection should be conducted and your evidence included with your report. Refer to the Compliance Program 7371.006, “Illegal Residues in Meat, Poultry, Seafood and other Animal Derived Foods” for in depth instructions on how to conduct a tissue residue inspection.

For information on tissue residue violations and activities you should contact the CVM/Division of Compliance (HFV-230, 240-276-9200).

5.9.6 - VETERINARY DEVICES

Medical devices for animal/veterinary use are not subject to the premarket approval requirements like human medical devices. Once an animal use device is marketed the Center is concerned with safety and efficacy of the veterinary device. CVM often recommends firms use the human device GMPs in controlling the manufacturing of animal use devices. CVM also suggests labeling be sent in for review by the Division of Compliance (HFV-230) to avoid misbranding. Regulatory questions for veterinary/animal use devices should be directed to the CVM/Division of Compliance (HFV-230).

5.9.7 - ANIMAL GROOMING AIDS

Grooming aids for animals formulated and labeled only to cleanse or beautify the animal are not cosmetics within the meaning of Section 201(i) and not subject to the Federal Food, Drug, and Cosmetic Act. Where animal grooming aids are labeled to contain an active drug ingredient or otherwise suggest or imply therapeutic benefit, they may be considered to be drugs and/or new animal drugs as defined by Section 201(v) of the Act (see CPG 653.100).

Questions on labeling and regulatory concerns should be directed to the Division of Compliance (HFV-230) at 240-276-9200.

5.9.8 - CVM BIO-RESEARCH MONITORING

Inspectional activities in the CVM bioresearch monitoring (BIMO) programs involve all animal product areas that have efficacy or safety requirements in order to be legally marketed. This includes Good Laboratory Practice (GLP) for Non-Clinical Laboratories; Sponsors, Monitors, Contract Research Organizations; and Clinical Investigators (CI). In most cases, inspections conducted under these programs will be done on assignment from CVM and occasionally with the participation of Center personnel as part of the inspection team.

SUBCHAPTER 5.10 – REPORTING

Following an inspection, you are required to prepare a report of your findings. Reporting includes the data and summary entered using FACTS, exhibits and attachments collected as evidence or for informational purposes and a
narrative report. Your narrative report should be prepared to accurately and concisely communicate the findings of your inspection and be adequate for its intended use. For example, an inspection of a new firm, one that FDA has not inspected previously, should be a comprehensive inspection focused on assessing the firm's compliance with applicable regulations. The resulting report would detail the products manufactured, the processes used to manufacture those products, the conditions of the environment in which products are manufactured or stored, any violations observed, persons responsible for the firm's operations, their actual duties and their responsibility for observed violations, distribution practices, and so on, providing information responsive to each of the required elements.

For establishments that have been previously inspected, you should determine what changes in operations and responsible individuals have occurred since the previous inspection, detail those changes in the narrative report and report on the areas of concern for the current inspectional outcome. For example, a non-violative inspection may only require a Summary of Findings report with the information required in the Summary, Administrative Data, General Discussion with Management, Voluntary Corrections, Refusals, Samples Collected, Exhibits Collected and Attachments (see IOM 5.10.4.1).

An OAI follow-up inspection that reveals continuing violations supporting a regulatory action would require the Summary, Administrative Data, Individual Responsibility and Persons Interviewed, Objectionable Conditions and Management's Response, Supporting Evidence and Relevance, Discussion with Management, Exhibits Collected, Attachments, and if appropriate, Refusals, Samples Collected, and Voluntary Corrections. Additionally, any information related to changes in previous operations would also need to be included in this type of report.

The key for you to remember in writing your narrative report is to communicate the findings of your inspection so that others may take the appropriate action. Notice that the required elements always include the product, interstate commerce, the violations observed and responsibility of firm officials. This is to document the elements of proof – Jurisdiction, Interstate Commerce, Violation and Responsibility (JIVR). Write your EIR with the intended use in mind. Your report may be a brief summary of an inspection of a firm in a state of compliance with applicable regulations all the way to a firm where the agency must take regulatory action to correct deficiencies.

5.10.1 - ESTABLISHMENT INSPECTION REPORT (EIR)

See IOM 1.1 English language requirement. The EIR consists of the following in this order: a printed copy of the FACTS Establishment Inspection Record (EI Record) including, at least, the endorsement with the EIR distribution printed at the bottom of the "endorsement" section of the EI Record; carbon or other copies of FDA forms issued during the inspection such as the FDA 482, FDA 483, and FDA 484; investigator's narrative report; copy of assignment if available; exhibits; and/or any additional material attached and referred to in the narrative report. Regarding the use of checklists that are completed during the inspection (such as the BSE Checklist), the original checklist should be submitted with the EIR. If you maintain the data in your regulatory notes, instead of entering the data directly on the checklist during the inspection, then a copy of the checklist that was completed using the data from your regulatory notes should be printed from FACTS and included with the EIR.

The signed original report is maintained in the district office or in the case of foreign inspections in the appropriate Center office. No copies of inspection reports will be maintained other than in the district and resident post files.

5.10.2 – ENDORSEMENT

The endorsement of the establishment inspection is prepared by the supervisor. Some supervisors may have the investigator prepare proposed endorsements. Endorsements should fit in the available space provided in FACTS. If the endorsement exceeds the 2000 character space provided in FACTS, a separate endorsement should be prepared, fully identifying the firm with a Summary of the Endorsement included in FACTS. The FACTS EI Record will be used as the endorsement and routing document to accompany the EIR. See also IOM 5.10.4.1.

Normally the endorsement consists of:

1. The reason for the EI, i.e., workplan, or assignments from headquarters. State the subject of the assignment and reference. If the assignment was issued hard copy (i.e. not through FACTS), it should be attached to the EIR following the narrative.
2. A brief history of previous findings including classification of previous EI, any action taken by the district and/or corrective action taken by the firm in response to inspectional observations from the previous inspection.
3. A concise summary and evaluation of current findings and samples collected.
4. Refusals, voluntary corrections or promises made by the firm's management.
5. Classification and follow-up consistent with inspectional findings and Agency policy including notification of other districts and headquarters as warranted.
6. Distribution consistent with District policy and the requirements of the specific Compliance Program and requirements as noted in IOM 1.7.3.

Note: Route a copy of the FACTS Establishment Inspection Record and the EIR to Division of Import Operations (DIO) when any violative, imported products
are identified. Per CPG 110.300, do not report the FURLS Registration number.

The existence of Personal Safety Alerts (IOM 5.2.1.3) or Personal Safety Plans (IOM 5.2.1.4) pertaining to the firm should be included in the endorsement section only and not in the EIR.

The signed endorsement should be updated to indicate if an addendum to the EIR (IOM 5.10.6) or an amended FDA 483 (IOM 5.2.3.1.6.1 and 5.2.3.1.6.2) has occurred.

**PROFILES:** Updating the Field Accomplishments and Compliance Tracking System (FACTS) database with a Compliance Status for each profile class code associated with the firm’s operations and/or products, is the responsibility of ORA Field and Center Investigators, Supervisors and Compliance Officers.

For Domestic inspections, hardcopy or e-mail notification of Potential OAI is not necessary. FACTS automatically sends OAI Notifications to OEIO/DCS electronically.

For foreign inspections, when a potential OAI Notification cannot immediately be entered in the FACTS firm profile record, the investigator should notify the Division of Food and Feed Program Operations and Inspections (CFSAN or CVM products) or the Division of Medical Products and Tobacco Inspections (CDER, CBER, CDRH, or CTP products) of the potential OAI situation via FAX (301-827-9791) as soon as the potential OAI situation is known and during the investigation. DFFPOI or DMPTI will then notify the appropriate Center.

See Exhibit 5-14 for more information on profiling CGMP/QS Compliance Status.

### 5.10.2.1 - Compliance Achievement Reporting System (CARS)

FACTS is used to report achieved and verified compliance actions, which are not the result of a legal action. A compliance achievement is the observed repair, modification, or adjustment of a violative condition, or the repair, modification, adjustment, relabeling, or destruction of a violative product when either the product or condition does not comply with the Acts enforced by the FDA. All CSOs should enter corrective actions into the CARS system as directed in 5.10.2.1.1. Each Supervisory CSO should verify that the corrections were entered by their CSO or should enter the information themselves.

#### 5.10.2.1.1 - REPORTING CRITERIA

There are three criteria for reporting into the CARS system:

1. **The detection or identification of the problem.** The problem may be observed by FDA, other federal officials, or by state or local authorities and referred to FDA; and as a result of an inspection, investigation, sample analysis, or detention accomplished by ORA or state officials under contract to ORA.

2. **The correction of the problem.** The correction is directly attributable to the efforts of ORA or state officials under contract to ORA (involving contract products only); and is unrelated to the filing of a legal action, i.e., seizure, prosecution, injunction.

3. **The verification of the correction of the problem.** The correction is verified by the FDA, other federal officials or state or local authorities and reported in writing to the FDA; and is based on an inspection, investigation, sample analysis, or letter from a firm to FDA certifying the problem has been corrected.

#### 5.10.2.1.2 - DATA ELEMENTS

Only when the corrective action(s) has been verified should a CARS be reported. The data elements are those entered/coded in FACTS (See IOM Exhibit 5-15):

1. **PAC.** See the Data Codes Manual. Should there be insufficient space to code all corrections verified on an occasion, record the most significant corrections.

2. **PROBLEM TYPE.** The problem type is the problem(s) identified during the operation(s). Use the List of Values (LOV) found in this field on the Compliance Achievement Reporting Screen. If “Other” is chosen, you should include an explanation in the “Remarks” field.

3. **CORRECTIVE ACTION.** The action the establishment took to correct the identified problem. Use the LOVs found in this field on the CARS screen. If “Other” is selected, you should include an explanation in the “Remarks” field.

4. **VERIFICATION DATE.** Use the date the corrective action(s) is verified, either through an establishment inspection, an investigation, or a letter from the establishment certifying the corrections have been made. Include documentation to verify the action such as repair receipts/plans.

5. **CORRECTING ORGANIZATION.** The FDA, other federal agency, or state or local authority, which observed the verified correction. Use the LOVs found in this field on the CARS screen.

6. **REPORTING ORGANIZATION.** The FDA, other federal agency, or state or local authority, which is actually inputting the verified correction. Use the LOVs found in this field on the CARS screen.

7. **REASON FOR CORRECTION.** The action the FDA took to make the correction happen. Use the LOVs found in this field on the CARS screen. If “Other” is chosen, you should include an explanation in the “Remarks” field.

### 5.10.3 - FACTS ESTABLISHMENT INSPECTION RECORD (EI RECORD)

Per FMD-130, each ORA District is responsible to ensure all investigators verify, correct, and enter changes to the OEI (including Profile data for profilable firms) on the firm’s maintenance screens in FACTS during each inspection, investigation and during any OEI update. Consult with your supervisor and District OEI Coordinator to assure data is accurately updated. See IOM Exhibit 5-16. The
FACTS generated assignment and FACTS Profile Data instructions are attached as IOM Exhibits 5-9 and 5-14.

Inspectional accountable time in FACTS consists of the hours devoted to file reviews (operational preparation), actual inspectional, investigational, time (onsite), document preparation (exhibit) and EIR (report) write-up. Accountable time does not include travel time. One occasional exception could be when more than one participant in an inspection/investigation travel together and discuss/prepare while in route.

5.10.3.1 – Inspection Basis

Compliance - Inspection is conducted to investigate potential violations that have not already resulted in an official agency action. These may include complaints (trade or consumer) which are not the primary reason for the inspection (otherwise see Consumer Complaint), recalls not classified as Class I, MedWatch Reports, Adverse Drug Experience Reports, information from confidential informants, etc.

Consumer Complaint - Inspection is conducted in direct follow-up to a consumer complaint. When a consumer complaint is received and the follow-up action chosen is to conduct an inspection to confirm allegations within the complaint or root causes that may have led to the condition described in the complaint, this value should be selected.

F/U to Class I Recall - Inspection is conducted in response to a Class I Recall conducted by the establishment. The inspection is conducted to determine the root cause and corrective actions addressing the violation(s) associated with the product.

F/U to Class I Recall and F/U to Injunction - Inspection is conducted in response to a Class I Recall conducted by the establishment AND pursuant to Permanent Injunction in accordance with the Consent Decree. In this instance, a firm under permanent injunction has conducted a Class I Recall and the inspection is conducted to determine the root cause and corrective actions addressing the violation(s) associated with the product. Additionally, the inspection covers the requirements of the Consent Decree for the Injunction.

F/U to Class I Recall and F/U to Warning Letter - Inspection is conducted in response to a Class I Recall conducted by the establishment AND f/u issues cited in a Warning Letter issued to the establishment. In this instance, a firm that has received a Warning Letter has conducted a Class I Recall and the inspection is conducted to determine the root cause and corrective actions addressing the violation(s) associated with the product in addition to covering corrective actions responsive to the violations cited in the Warning Letter. The Warning Letter may have issued as a result of the previous inspection or other circumstance.

F/U to Class I Recall and OAI Inspection F/U - Inspection is conducted in response to a Class I Recall conducted by the establishment AND f/u to previous OAI-classified inspection, where a regulatory or administrative action has not been completed. This value captures the situation where the previous inspection of the firm was classified OAI, but no official action was taken and the firm has conducted a Class I Recall. The inspection is focused on the root causes of the violations leading to the recall and may also address previously cited violations. Before conducting an inspection of a firm where the previous inspection was classified OAI with no regulatory action taken, be sure to discuss what areas to cover with your supervisor and/or compliance officer.

F/U to Injunction - Inspection is conducted pursuant to Permanent Injunction and in accordance with the Consent Decree.

F/U to Warning Letter - Inspection is conducted to follow-up issues cited in a Warning Letter issued to the establishment.

OAI Inspection F/U - Inspection is conducted to follow-up previous OAI-classified inspection where a regulatory or administrative action has not been completed. There can be a number of situations where an action is not taken although the observations cited during the previous inspection met the threshold for an OAI classification. Consult your supervisor and/or compliance officer prior to initiating these types of inspections.

Surveillance - Inspection is conducted as a routine assignment with no other indicators of non-compliance. For example, an inspection of a firm whose previous inspection was classified NAI; there have not been any complaints or recalls, etc.

5.10.4 - NARRATIVE REPORT

See IOM 1.1, English language requirement. You should use Turbo EIR for all EIRs. The narrative report is the written portion of the EIR, which accurately describes the investigator’s inspectional findings. The narrative report may be prepared in two formats depending on the type of inspection and inspection classification. A Summary of Findings narrative report may be used for non-violative, non-initial inspections - see IOM 5.10.4.1. The full Standard narrative report is used for initial and potential Official Action Indicated (OAI) classified inspections - see IOM 5.10.4.3. The "Summary of Findings" report format may be used for some Voluntary Action Indicated (VAI) classified inspections as directed by your supervisor. Additional requirements for human drug and medical device reports are described in IOM 5.5.8 and 5.6.9. For all reporting formats, include additional information as directed by your assignment, Compliance Program Guidance Manual, or your Supervisor.

All reports should be prepared as stand-alone documents outside of FACTS. Your Establishment Inspection Report (EIR) should:
1. Be factual, objective, and free of unsupportable conclusions.
2. Be concise and descriptive while covering the necessary aspects of the inspection.
3. Not include opinions about administrative or regulatory follow-up.
4. Generally, be written in the first person using the active voice.
5. Be signed by all FDA and commissioned personnel participating in the inspection. See IOM section 5.1.2.5.1 when more than one FDA or commissioned person participated in the inspection.

Refer to IOM 5.10.6 for an Addendum to EIR.

**5.10.4.1 - Non-Violative Establishments**

Investigators should use "Summary of Findings", stand-alone, narrative reports for non-violative domestic establishments, unless otherwise directed by your supervisor, the assignment or the Compliance Program Guidance Manual.

The Summary of Findings Report may not be written solely in the FACTS provided "Inspection Summary" heading. The Summary of Findings report should include:

1. The reason for the inspection;
2. The date, classification and findings of the previous inspection;
3. The actual inclusive dates of the inspection (these may be included as part of a header or in the body of the EIR.)
4. The name of the person to whom credentials were shown and the FDA 482, Notice of Inspection was issued and the person's authority to receive the FDA 482. Explain if you were unable to show credentials or issue forms to top management. Include the name of the person to whom FMD-145 correspondence should be directed;
5. The scope of the inspection; i.e., comprehensive or directed; and a brief description of the products, processes or systems covered during the inspection; the manufacturing codes and if necessary their interpretation.
6. Significant changes (e.g., personnel, facilities, products, processes) since the previous inspection
7. The significant findings if any;
8. Management's response or corrections;
9. Warnings given to management; and
10. The investigator's handwritten signature.

**5.10.4.2 - Violative Establishments**

For domestic inspections where regulatory action is being recommended and when the district has final classification responsibility, the inspection report should normally be submitted within 10 days to the District or Center Compliance Branch as per established procedures. Please note, that depending on the type and severity of the regulatory action, it may be necessary to submit the EIR in less than 10 days. You should consult with your supervisory investigator in these instances. Refer to FMD-86 and the RPM regarding other timeframes associated with non-violative inspections.

All violative EIR's should in addition to the information required for non-violative reports contain the following:

1. The objectionable conditions or practices described in sufficient detail so someone reading the report will clearly understand the observation(s) and significance.
2. The objectionable conditions or practices cross-referenced to FDA 483 citations, samples collected, photographs, or other documentation including exhibits attached to the EIR.
3. Information as to when the objectionable conditions or practices occurred, why they occurred, and who is or was responsible, developed to the highest level in the firm.

**5.10.4.3 - Individual Narrative Headings**

There are many acceptable ways of organizing a narrative report. The key is to cover the required information in IOM 5.10.4 and 5.10.4.2, or as required by the assignment, Compliance Program Guidance Manual, or your supervisor. The appropriate use of headings should not result in repetition of the same information in different sections. You are encouraged to create headings as necessary to present the inspectional findings in the most concise manner. For non-violative, a single heading such as "Summary of Findings" is sufficient (for exceptions, see IOM 5.10.4.1). Turbo EIR should be used to generate the FDA 483. In certain instances, if you experience computer problems, do not delay the issuance of the FDA 483. See IOM 5.2.3. You should use Turbo EIR for all EIRs.

**5.10.4.3.1 - STANDARD NARRATIVE REPORT**

This is intended to outline the minimal information needed to produce a narrative report that supports further agency regulatory action, as warranted. Investigators are encouraged to add additional report headings as needed to communicate important information about the inspection, relevance of inspectional observations that may impact public health, and /or to address specific requests from directed assignments.

**Comprehensive Reports (Include all applicable sections)**

A comprehensive EIR should be prepared for initial inspections in all program areas. It is essential to describe the products manufactured, the process the manufacturing and storage environment, distribution patterns/interstate commerce, individual responsibility of key employees, history of business, all objectionable conditions observed, etc. All things pertinent to the operations and management of the establishment should be included in these reports. The comprehensive report may also be used for other situations requiring full reporting such as Routine Surveillance - OAI. An abbreviated inspection does not necessarily equate to an abbreviated report.

Required elements

- 5.10.4.3.2 – Summary
• 5.10.4.3.3 – Administrative data
• 5.10.4.3.4 – History
• 5.10.4.3.5 – Interstate (I.S.) Commerce
• 5.10.4.3.6 – Jurisdiction (Products Manufactured and/or Distributed)
• 5.10.4.3.7 – Individual Responsibility and Persons Interviewed
• 5.10.4.3.8 – Firm’s Training Program
• 5.10.4.3.9 – Manufacturing/Design Operations
• 5.10.4.3.10 – Manufacturing Codes

Routine Surveillance – NAI Reports

When FDA has an inspectional history for the firm and no deficiencies were observed by the investigator, a brief report may be prepared. The intent of this report is to include only the required information about the firm and what areas were covered during the inspection. “Change reporting” means information that differs from the previous inspection report such as changes in management, products produced, manufacturing processes, etc. Where these changes have occurred, the applicable section heading in the EIR should be included. The elements may also be captured in a “Summary of Findings Only” report without header information. See IOM 5.10.4.1.

Required elements

• 5.10.4.3.2 – Summary
• 5.10.4.3.3 – Administrative data
• 5.10.4.3.14 – Refusals
• 5.10.4.3.15 – General Discussion with Management
• 5.10.4.3.16 – Additional Information
• 5.10.4.3.17 – Samples Collected
• 5.10.4.3.18 – Voluntary Corrections
• 5.10.4.3.19 – Exhibits Collected
• 5.10.4.3.20 – Attachments

Change reporting only

• 5.10.4.3.4 – History
• 5.10.4.3.6 – Jurisdiction (Products Manufactured and/or Distributed)
• 5.10.4.3.7 – Individual Responsibility and Persons Interviewed

Routine Surveillance – VAI Reports

For firms with an inspectional history and the outcome of the inspection is a VAI classification, the below elements would be required, plus change reporting. Note that the difference in the NAI versus the VAI report is the inclusion of narrative addressing objectionable conditions observed during the inspection. Each objectionable condition or practice must be documented in the EIR along with discussion of the evidence, relevance and discussion with management.

Required elements

• 5.10.4.3.2 – Summary
• 5.10.4.3.3 – Administrative data
• 5.10.4.3.14 – Refusals
• 5.10.4.3.15 – General Discussion with Management
• 5.10.4.3.16 – Additional Information
• 5.10.4.3.17 – Samples Collected
• 5.10.4.3.18 – Voluntary Corrections
• 5.10.4.3.19 – Exhibits Collected
• 5.10.4.3.20 – Attachments

Change reporting only

• 5.10.4.3.4 – History
• 5.10.4.3.7 – Individual Responsibility and Persons Interviewed
• 5.10.4.3.8 – Firm’s Training Program
• 5.10.4.3.9 – Manufacturing/Design Operations
• 5.10.4.3.10 – Manufacturing Codes

Routine Surveillance – OAI

For an OAI surveillance inspection, follow the guidance under Comprehensive Reports.

OAI Follow-up Inspection Reports

OAI follow-up inspections are inspections conducted following an OAI classified inspection. Inspections of this nature are conducted to determine whether corrective actions have been implemented or significant violations continue. The outcome of these inspections may range from NAI to OAI. The intended use of the EIR should be the driving force of the content of these EIRs. Typically, the follow-up should be done relatively soon after the previous inspection, so changes to products, process, personnel, etc. should be minimal. The NAI and VAI reports should focus on corrective actions implemented by firm management to correct the violative conditions observed during the previous OAI inspection. Those
reports may be Summary of Findings only or may follow
the other NAI and VAI report formats above. An OAI
follow-up inspection may lead to a regulatory action such
as seizure, injunction and/or prosecution. Those reports
should focus on documenting the continuing violations,
responsibility for those violations, any corrective actions
implemented or inadequate corrective actions and defining
the new scope of violations observed including the
products affected. Scope should include additional lots,
products, timeframe and distribution. These reports should
document all elements of JIVR (Jurisdiction, Interstate
Commerce, Violation and Responsibility). This allows for
the report to support whatever regulatory action is deemed
necessary.

Required elements

- 5.10.4.3.2 – Summary
- 5.10.4.3.3 – Administrative data
- 5.10.4.3.7 – Individual Responsibility and Persons
  Interviewed
- 5.10.4.3.13 – Objectionable Conditions and
  Management’s Response
- 5.10.4.3.13.1 – Supporting Evidence and
  Relevance
- 5.10.4.3.13.2 - Discussion with Management
- 5.10.4.3.14 – Refusals
- 5.10.4.3.17 – Samples Collected
- 5.10.4.3.18 – Voluntary Corrections
- 5.10.4.3.19 – Exhibits Collected
- 5.10.4.3.20 – Attachments

Change Reporting Only:

- 5.10.4.3.5 – Interstate (I.S.) Commerce
- 5.10.4.3.6 – Jurisdiction (Products Manufactured
  and/or Distributed)
- 5.10.4.3.9 – Manufacturing/Design Operations
- 5.10.4.3.10 – Manufacturing Codes
- 5.10.4.3.11 – Complaints

5.10.4.3.2 - SUMMARY

1. Provide the reason for the inspection (e.g., compliance
   program, by assignment, etc.);
2. The scope of the inspection (comprehensive, directed,
   sample collection only, QSIT level, etc.);
3. Provide a summary of the findings, date, and
classification of the previous inspection and the firm’s
response/corrective actions.
4. List the products, systems and processes covered
during the current inspection, and the types of records
and documents reviewed. For human drug reports, list
the systems not covered.
5. Provide a summary of the current findings, refusals,
samples collected, warnings given to management,
and a summary of management’s response or
voluntary corrections.
6. Per CPG 110.300, do not report the FURLS
   Registration number.

5.10.4.3.3 - ADMINISTRATIVE DATA

Administrative Data:
1. The firm name, address, phone, FAX and e-mail
   address.
2. Report the names and titles of the Investigator(s),
   Analyst(s), non-FDA officials, etc. Report the name of
   the firm’s responsible official who gave permission to
   non-FDA officials without inspection authority to
   accompany you during your inspection. See IOM 5.1.1
   and 5.2.2.
3. The inclusive date(s) of the current inspection, i.e., list
   the actual dates in the plant.
4. If a team inspection and some individuals were not
   present during the entire inspection, indicate dates in
   plant for each team member.
5. For foreign inspections with Locally Engaged Staff
   (LES)/Foreign Service National (FSN) participation
   include this language:

This inspection was supported by
____________________ (during the period of
__________), who is a Locally Engaged Staff (LES)
hired by the United States Embassy and assigned to
FDA to work in support of FDA activities. All
information, including documents collected during this
inspection and any translation from local language to
English by ______________(LES) that supports the
Form FDA 483, Inspectional Observations (if Form
FDA 483 was issued) and the Establishment Inspection
Report (EIR) was collected in collaboration with the
FDA investigator(s).

Report Full Names and Titles of:
1. To whom FDA Official Credentials were shown,
2. To whom any FDA forms were issued to or signed by
during the inspection (FDA 482, 483, 484, 463, etc.);
   where appropriate, explain the reason a form(s) was
   not issued to or signed by the most responsible indi
   vidual (this may be reported in the Individual Respon­
   sibility and Persons Interviewed heading below),
3. Who wrote which section of the EIR, if this was a team
   inspection report,
4. In-plant inspectors or other government agencies (IOM
   5.4.9), and
5. For domestic and foreign food facilities, document to
   whom the FSMA Fee Information Sheet was provided
to.

5.10.4.3.4 - HISTORY

History:
1. Report the legal status of the firm (corporation, partner­
   ship, limited liability company, etc.). If a corporation, list
   in which state and when the firm was incorporated.
2. List the parent corporation, corporate address and any
   subsidiaries.
3. Provide a summary of any regulatory actions and prior
   warnings (do not cite any action only recommended
but not approved). You should also report any significant/relevant inspectional history pertinent to the current EI or recommendation.
4. Include any relevant recalls, etc. since the last inspection.
5. Report the hours of operation and any changes from past inspections (include seasonal variations).
6. Report the current registration(s) status or any changes to registration status. Per CPG section 110.300, do not report the FURLS Registration number.
7. If directions to the firm would be helpful in future visits, include the information.
8. Provide the names, titles and addresses of top management official(s) to whom correspondence should be addressed (FMD 145, W/L, etc.).
9. For foreign inspections, list U.S. consignees to whom the firm’s products are shipped.
10. For Human Drugs - domestic firms, identify the general types of customers and provide the names and addresses for several regular customers of a few of the firm’s products.

5.10.4.3.5 - INTERSTATE (I.S.) COMMERCE

Interstate (I.S.) Commerce:
1. Report changes in the previous estimate of the percentage of products shipped outside of the state (or exported to the U.S.) and the basis of the estimate.
2. Report the firm’s general promotion and distribution patterns.
3. If there is an apparent violative product, provide examples of I.S. shipments of violative product(s); or
4. If no such shipments, provide examples of I.S. shipments of major components of apparent violative products - with complete I.S. documentation in either case.

5.10.4.3.6 - JURISDICTION (PRODUCTS MANUFACTURED AND/OR DISTRIBUTED)

Jurisdiction (Products Manufactured and/or Distributed):
1. Include a list of a representative number of currently marketed products subject to FD&C Act or other statute enforced by FDA or counterpart state agency, including any believed violative.
2. Collect appropriate labeling (product and case labels, inserts, brochures, manuals, promotional materials of any type) for those products believed violative or representing any significant new or unusual operation, industry or technology; or as directed by your supervisor.
3. Document any applicable labeling agreements (and obtain a copy) and statutory guaranty given or received per Sections 301(h) and 303(c)(2) of the FD&C Act [21 U.S.C. 321(h) and 333(c)(2)] (IOM 5.3.7.2)

In addition, the label, labeling and promotional materials are a critical part of determining a product’s intended use.
1. In instances where a regulatory action is being considered based on product labels, labeling, and/or other promotional materials, including any Internet websites, you should collect all available documentation. This includes all written, printed or graphic matter on the immediate container of an article or accompanying the article (the product’s label and labeling, see FD&C Act, 201(k) and (m) [21 U.S.C. 321(k) and (m)] and IOM 4.4.9.1). Accompanying labeling could include brochures, pamphlets, circulars, and flyers, as well as audio and video tapes.

2. In cases where there may be a dispute about whether a product is a drug or a dietary supplement, you should collect all materials which claim a product can be used for the treatment of any disease.

5.10.4.3.7 - INDIVIDUAL RESPONSIBILITY AND PERSONS INTERVIEWED

Report with whom you dealt, and in what regard (both during and prior to the start of the inspection):
1. Who provided relevant information,
2. Who accompanied you during the inspection,
3. Who refused access to required records or any other refusal of information (Note: a separate heading for Refusals may be needed if refusals are significant, extensive or an Inspection Warrant is anticipated),
4. Who refused to permit inspection (IOM 5.2.5.1) and
5. For Human Drug inspection reports, also include the name, title, physical mailing address, phone, and fax number and e-mail address for any U.S. Agent or broker who represents the company when dealing with the FDA.

Describe roles and authorities of responsible individuals, including the full names and titles of individuals providing you with information.

Describe roles, authorities and responsibilities of officials at headquarter or corporate organizations for this firm; including their names, titles and addresses.

Report changes to the following:
1. Who is the most responsible individual at the inspected firm? Who is the responsible head or designated correspondent? Refer to IOM 5.3.6, 5.3.6.1, and 5.3.6.2.
2. Report full names and titles of owners, partners, and corporate officers. Who has the duty, power and responsibility, and authority to prevent, detect, and correct violation(s), and how is this demonstrated and/or documented? See IOM 5.3.6.2.
3. Report the chain of command; include an organizational chart (create if necessary).
4. Obtain a copy of public annual report, if any.
5. List the names and titles of key operating personnel.

5.10.4.3.8 - FIRM’S TRAINING PROGRAM

The firm’s training programs are of particular significance where inspectional findings find people may not be adequately trained.

5.10.4.3.9 - MANUFACTURING/DISIGN OPERATIONS

Manufacturing/Design Operations:
1. Report only changes to the firm’s general overall operations, including significant changes in equipment,
processes, or products since the previous inspection. Include schematics, flow plans, photographs, formulations and diagrams, if useful.

2. List names and sources of new or unusual components or raw materials.

3. Report equipment considered new or unusual unless otherwise directed.

4. Submit pertinent formulas (especially those being manufactured during your inspection) and processing instructions with labeling of suspect products.

For human drug inspection reports:

This section of the EIR should be organized by system covered during the EI as outlined in CP 7356.002. In each section, include a brief summary of what you reviewed in order to meet the key system element outlined in the CP. You should add more detail for the system elements found to be deficient, or the subject of a FDA 483 observation.

For medical device inspection reports:

1. Describe manufacturing operations by sub system covered in your inspection (Management Controls, Design Controls, Production and Process Controls, Corrective and Preventive Action Controls, Material Controls, Facility and Equipment Controls, and Records/Documents/Change Controls). For ALL Level 2, 3, and "for cause" inspections: for production and process controls - indicate which production processes were covered/reviewed. If a subsystem was not specifically covered during your EI, you do not need to separately describe the general operations of that subsystem.

2. For all inspections covering CAPA - indicate which data sources were available for review and which were actually reviewed; include a brief statement regarding coverage or non-coverage of applicable tracking requirements, MDRs, sterilization, and reports of corrections and removals.

3. If the Design Control system was covered, indicate the design project(s) covered during the inspection. Where design activities occur at a location other than the manufacturing site, list the name, address of the design location and responsibilities of those performing the design activities.

4. If applicable, identify the name and address of the specification developer if different from either the manufacturing site or where design activities occur.

5.10.4.3.10 - MANUFACTURING CODES

Manufacturing Codes

1. If the manufacturing codes are unchanged, include a statement in the EIR the system is the same as described in reports on file at the District. Indicate the date of the EIR in which the codes are fully explained.

2. If the manufacturing codes have changed, describe the manufacturing coding system (lot, batch, product, etc.), and a key to interpretation of codes.

3. For medical device inspections reports: where appropriate, include a description of the system used to identify and maintain control of components during the manufacturing process, as well as, the codes used for traceability (for applicable finished devices).

5.10.4.3.11 - COMPLAINTS

Note: These complaints include those reported to the FDA by consumers, health care professionals, industry, etc.; and all complaints received by the firm.

1. Report your review of the firm’s complaint file(s).

2. In addition, if returned goods and/or documents for returned goods are examined, describe findings. If not examined, so indicate.

3. Report your follow-up of consumer/trade complaints, Adverse Event Reports, MDR’s, MedWatch reports or recalls identified in the district factory jacket for coverage. Correlate consumer/trade complaints, Adverse Event Reports, MDR’s, MedWatch reports to specific objectionable conditions observed.

5.10.4.3.12 - RECALL PROCEDURES

Describe plans and procedures for removing products from marketing channels if necessary. If these procedures are in written SOP-type format, you may reference any copies obtained to aid in your explanation.

5.10.4.3.13 - OBJECTIONABLE CONDITIONS AND MANAGEMENT’S RESPONSE

If any observations were provided to management in writing (FDA 483) at the conclusion of the inspection list each observation and report each observation providing information organized under the two headings Supporting Evidence and Relevance, and Discussion with Management below.

NOTE: Observations of a verbal nature (i.e., Discussion Items) should be reported in sufficient detail under the General Discussion with Management (correlate any Exhibits, samples, etc. to any "verbal" observations).

5.10.4.3.13.1 - Supporting Evidence and Relevance

Sufficiently describe the observation as necessary to relate the facts as you found them.

1. Identify specific pages of exhibits and/or samples (e.g., procedure title, section, paragraph, sentence), labeling text, interstate shipping records which in your judgment document violations so supervisors, compliance officers, and other reviewers can readily evaluate your evidence.

2. Describe verbal statements (verbatim if possible) by firm officials having knowledge, duty, power, and responsibility to detect, prevent, or correct the apparent violation.

3. Identify the responsible party for each apparent violation (i.e., if known.)

4. Identify which team member (if applicable) was responsible for the observation.

5. When appropriate explain how this observation relates to the overall situation; i.e., impact on the product,
batches, or lots involved, and any relationship to other products, processes, or other FDA 483 observations.
6. The duration of the problem.

5.10.4.3.13.2 - Discussion with Management

Discussion with management:
1. Report management's response to each specific observation, time frames given for corrections and/or corrective action.
2. Report any disagreements with or refusals to correct the observation.

For medical device inspection reports:
1. For each observation based on sampling of records, indicate which Sample Table and level of confidence was used and the actual number of records sampled.
2. If the number sampled is different than the actual number reviewed, so indicate.

5.10.4.3.14 - REFUSALS

Provide full details of all refusals of/for requested information, statutory information, photography, entry, etc. received during the inspection, including who made the refusal and, if available, why the refusal was given.

In the case of drug inspections, similarly provide full details of all instances of delaying, denying, limiting, or refusing an inspection encountered during the inspection.

5.10.4.3.15 - GENERAL DISCUSSION WITH MANAGEMENT

General Discussion with Management:
1. Report the names and titles of all present, including those present via electronic media (describe).
2. Include the name and title to whom the FDA 483 was issued.
3. Provide additional discussion items not provided in writing at the conclusion of the inspection, such as: questionable labels, labeling and/or labeling practices, commercialization of products covered by IDE or IND, fraudulent health claims, registration/listing deviations, lack of approved PMA, 510(k), NDA, ANDA, etc. These include all verbal observations deemed not to merit inclusion on the FDA 483 (IOM 5.2.3)
4. A description of each warning, recommendation, or suggestion given to the firm, and to whom given.
5. Management's general responses to the inspection and/or to groups of items listed on the report of observations or discussed at the conclusion of the inspection.

5.10.4.3.16 - ADDITIONAL INFORMATION

Report changes as appropriate.
1. Describe contractors used and for what purpose. For Medical Device inspection reports: also include names and addresses of all applicable third party installers or servicing organizations used by the manufacturer. Include their responsibilities.
2. Describe suppliers (major raw material, active ingredient, etc.) used and for what.
3. During inspections, when violative products imported into the U.S. or intended to be imported into the U.S., are encountered, document the product and foreign manufacturer in the EIR. Violative products could be rejected APIs due to non-conformance with the USP, foods without appropriate labeling, etc. Send a copy of the EIR to OEIO/DIO. See IOM 5.2.1 and 5.10.2.
4. For initial inspections, verify distribution patterns for the firm's products, raw materials, and components to firms which warehouse or further process products which may be subject to FDA regulations. Districts should incorporate information obtained into their Official Establishment Inventory improvement activities and complete form FDA 457, Product/Establishment Surveillance Report as appropriate. See IOM 8.6.2.
5. Report pertinent facts, which do not fit another section of the EIR. (For firms located in foreign countries, include information relative to lodging and travel; for domestic firms, include information relative to location of firm if difficult to find; etc.).

For human drug inspection reports - PDMA Coverage:
1. Describe what sample loss, theft, or diversion reports were covered during the inspection.
2. Describe the firm's sample audit and security systems, including a review of the firm's SOP's. Significant problems which may contribute to the firm's inability to adequately monitor sample distribution via sales representative, mail or common carrier should be addressed under objectionable conditions.

5.10.4.3.17 - SAMPLES COLLECTED

List and describe samples collected during the inspection.

5.10.4.3.18 - VOLUNTARY CORRECTIONS

Voluntary Corrections:
1. Provide a brief description of improvements initiated by the firm in response to a previous inspection, report of observations and/or a warning letter.
2. Report voluntary destructions, recalls, and similar actions since the prior inspection or during this inspection.
3. Report any follow-up to recalls identified during the inspection (may be by referencing Attachment B recall report).
4. Include recalls to specific objectionable conditions observed.
5. Provide the identity of person(s) responsible for the corrections.
6. Report any appropriate voluntary corrections in FACTS CARS.

5.10.4.3.19 - EXHIBITS COLLECTED

List all exhibits attached. See IOM 5.10.5, Exhibits.
Briefly, describe or title each exhibit attached. You should include in your description the number of pages for each Exhibit listing.

NOTE: For complex inspections a cross-reference from the FDA 483 and verbal observations to applicable exhibits and samples can be useful during further review.

5.10.4.3.20 – ATTACHMENTS

Attachments as referred to here are any material not provided by the firm during the inspection and referred to in the EIR, which are not evidentiary in nature; such as assignments, Center provided protocols, website information printed during inspectional preparation, etc. Non-evidentiary material attached to the narrative portion of the EIR should be identified as “Attachments” similar to IOM 5.3.8.2. Documents attached to the EIR may be referred to under the attachments heading, such as a copy of the FDA 463a, the FDA 482, FDA 483, etc. (in form number order); but such documents/forms may not be numbered, altered from their issued state, bear adhesive identification labels, etc. See the opening sentence of IOM 5.10.5. List and attach copies of associated reports (Recall Attachment B Report, etc.).

5.10.4.3.21 - SIGNATURE

All participants will sign the final narrative portion of the EIR. The prescribed format is to include each person’s name and title. Participants should include their District and Resident Post (or other affiliation) below the signature if needed. In some cases immediate signature by all participants is not possible. An example as to how this can be accomplished is to forward an electronic “draft” copy of the EIR for all to read and approve, then followed or accompanied by the original signature sheet. When signed, return to the lead investigator for proper filing and routing. When using this method, a photocopy of the original signature page is made with the lead investigator’s signature and temporarily attached to the EIR.

5.10.5 - EXHIBITS

Exhibits are materials collected from the firm during the inspection and do not include FDA forms, copies of assignments, or information obtained outside of the firm. For example, website downloads printed prior to the start of the inspection are not exhibits. Collect only records and documents which are pertinent to the inspectional findings or are required by assignment or Compliance Program. Exhibits should contribute to the objective of the assignment and the clarity of the report. They may include flow-plans, schematics, layouts, etc. Additional exhibit examples include copies of procedures or batch records that relate to and provide evidence of a violation. The exhibits should be discussed and referenced in your narrative report and attached to the final Establishment Inspection Report. If the materials collected from the firm are not needed as exhibits, they should be destroyed in accordance with district policy. Copies of procedures or patient records that do not serve as evidence of a violation should not be collected unless you are directed to do so.

Exhibits which include medical records obtained during an investigation or inspection should be handled in accordance with current personal privacy disclosure rules. Such patient records should remain intact and stored in the official files. When copies of these records are requested internal to FDA, they should be redacted by obliterating the patients full name (keeping first and last initial only), social security number, date of birth, race, personal address and any other personal identifiers. All external requests should be handled by the FOI officer.

Submit at least three copies of new or suspect labeling or other material collected as exhibits for labeling purposes. See IOM 4.4.9 for exceptions. These should be mounted in a manner so complete sets are submitted that can be reviewed by individuals in separate offices, i.e., labels 1-10 in each of three sets. You should identify records/exhibits submitted with an EIR using at least the Exhibits’ number, firm name, date(s) of the inspection, and your initials. See IOM 5.3.8.2.

5.10.5.1 - Electronic Records as Exhibits

Electronic records included as exhibit to the EIR should be stored to protect the integrity of the data. Refer to IOM 5.3.8.3.3.1. Electronic records should be protected from degradation, including preventing exposure of the electronic storage media to extreme temperatures and magnetic fields where necessary. Additional precautions to preserve the electronic records may be required, and you should be guided by your District procedures for handling electronic storage media. See IOM 5.10.4.3.20 Attachments and 5.10.5 Exhibits.

5.10.6 - ADDENDUM TO EIR

If your EIR requires correcting or clarification after it has been endorsed, signed and distributed (outside of the District Office), an addendum may be prepared at the request of your supervisor.

The addendum will be written in Turbo EIR if the original EIR was written in that program (See 5.10.6.1). If the original EIR was written outside of Turbo EIR, the addendum will be written in Word and appended to the original EIR. The addendum should clearly identify itself with the EIR being added to, explain the necessity for the addendum, and clearly define what section(s) and page(s) are being revised. The addendum will be signed by the preparer(s) in a new signature block added to the end of the “Addendum” section. Sign only the “Addendum” page(s) for submission to your supervisor. The Addendum pages should be added to the original EIR you submitted (See 5.10.6.2 and Exhibit 5-19).

The addendum must be endorsed in FACTS by the supervisor (See 5.10.1).

5.10.6.1- How to Retrieve a Completed EIR from Turbo
Below are instructions to retrieve a completed EIR from the server to make changes to your EIR if it has already been set to Complete in Turbo (For those districts who use Turbo):

1. Click on the “File” menu in Turbo, while connected to the network.
2. Select “Retrieve a Specific Assignment from the Server.”
3. Select “Completed Assignment.”
4. Select “Yes” to acknowledge that you are retrieving a signed and completed document.
5. Select the desired EIR from the list of documents and click “OK”. Your EIR is now located in your Turbo inbox.
6. Expand the “Completed” folder from the Turbo folder list and select the desired EIR.
7. Click “Modify (Signed)” button and click “OK” to modify EIR.
8. Click the “Check Out” Button.
9. Click the “Edit/Print” button to open and create an addendum.

If you need additional assistance, please contact the Turbo Help Desk.

5.10.6.2 – Instructions on How to create the Addendum

1. Find the signature block located on the last page of the EIR.
2. Place the cursor below the signature block and insert a “section break → next page”. At this step, it is important to note that the body, header, and footers of the original EIR must not change. Therefore, the Addendum section must begin pagination as page 1 of X.
3. Add section heading on new page titled “ADDENDUM”.
4. In the body of the Addendum, explain the reason for the addendum and clearly define what section(s) and page(s) are being revised, or what information is being added or clarified.
5. Insert a new signature block at the end of the Addendum section.
6. Print and sign only the “Addendum” page(s) for submission to your supervisor.
7. Save the entire document.
Notice of Inspection is hereby given pursuant to Section 704(a)(1) of the Federal Food, Drug, and Cosmetics Act [21 U.S.C. 374(a)] and/or Part F or G, Title III of the Public Health Service Act [42 U.S.C. 262-264].

As a small business that is subject to FDA regulation, you have the right to seek assistance from the U.S. Small Business Administration (SBA). This assistance includes a mechanism to address the enforcement actions of Federal agencies. SBA has a National Ombudsman’s Office that receives comments from small businesses about Federal agency enforcement actions. If you wish to comment on the enforcement actions of FDA, CALL (888) 734-3247. The website address is www.sba.gov/ombudsman.

FDA has an Office of the Ombudsman that can directly assist small business with complaints or disputes about actions of the FDA. That office can be reached by calling (301) 796-8530 or by email at ombuds@oc.fda.gov.

For industry information, go to www.fda.gov/ooc/industry.

1 Applicable portions of Section 704 and other Sections of the Federal Food, Drug, and Cosmetic Act [21 U.S.C. 374] are quoted below:

Sec. 704(a)(1) For purposes of enforcement of this Act, officers or employees duly designated by the Secretary, upon presenting appropriate credentials and a written notice to the owner, operator, or agent in charge, are authorized (A) to enter, at reasonable times, any factory, warehouse, or establishment in which food, drugs, devices, tobacco products, or cosmetics are manufactured, processed, packed, or held, for introduction into interstate commerce or after such introduction, or to enter any vehicle being used to transport or hold such food, drugs, devices, tobacco products, or cosmetics in interstate commerce; and (B) to inspect, at reasonable times and within reasonable limits and in a reasonable manner, such factory, warehouse, establishment, or vehicle and all pertinent equipment, finished and unfinished materials, containers, and labeling therein. In the case of any person (excluding farms and restaurants) who manufactures, processes, packs, transports, distributes, holds, or imports foods, the inspection shall extend to all records and other information described in section 414, when the standard for records inspection under paragraph (1) or (2) of section 414(a) applies, subject to the limitations established in section 414(d). In the case of any factory, warehouse, establishment, or consulting laboratory in which prescription drugs, nonprescription drugs intended for human use, restricted devices, or tobacco products are manufactured, processed, packed, or held, inspection shall extend to all things therein (including records, files, papers, processes, controls, and facilities) bearing on whether prescription drugs, nonprescription drugs intended for human use, restricted devices, or tobacco products which are adulterated or misbranded within the meaning of this Act, or which may not be manufactured, introduced into interstate commerce, or sold, or offered for sale by reason of any provision of this Act, have been or are being manufactured, processed, packed, transported, or held in any such place, or otherwise bearing on violation of this Act. No inspection authorized by the preceding sentence or by paragraph (3) shall extend to financial data, sales data other than shipment data, pricing data, personnel data (other than data as to qualifications of technical and professional personnel) performing functions subject to this act (Continued on Reverse)
Act), and research data (other than data relating to new drugs, antibiotic drugs, devices, and tobacco products and subject to reporting and inspection under regulations lawfully issued pursuant to section 505(i) or (k), section 519, section 520(g), or chapter IX and data relating to other drugs, devices, or tobacco products, which in the case of a new drug would be subject to reporting or inspection under lawful regulations issued pursuant to section 505(j). A separate notice shall be given for each such inspection, but a notice shall not be required for each entry made during the period covered by the inspection. Each such inspection shall be commenced and completed with reasonable promptness.

Sec. 704(a)(2) The provisions of the third sentence of paragraph (1) shall not apply to (A) pharmacies which maintain establishments in conformance with any applicable local laws regulating the practice of pharmacy and medicine and which are regularly engaged in dispensing prescription drugs or devices, upon prescriptions of practitioners licensed to administer such drugs or devices to patients under the care of such practitioners; or (B) upon prescriptions of practitioners licensed to administer such drugs or devices to patients under the care of such practitioners in the course of their professional practice, and which do not, either through a subsidiary or otherwise, manufacture, prepare, propagate, compound, or process drugs or devices for sale, other than in the regular course of their business; or (C) practitioners licensed by law to prescribe or administer drugs or prescribe or use devices, as the case may be, and who manufacture, prepare, propagate, compound, process drugs, or manufacture or process devices solely for use in the course of their professional practice; (D) persons who manufacture, prepare, propagate, compound, or process drugs, or manufacture or process devices solely for use in research, teaching, or chemical analysis and not for sale; (E) such other classes of persons as the Secretary may by regulation exempt from the application of this section upon a finding that inspection as applied to such classes of persons in accordance with this section is not necessary for the protection of the public health.

Sec. 704(a)(3) An officer or employee making an inspection under paragraph (1) for purposes of enforcing the requirements of section 412 applicable to infant formulas shall be permitted, at all reasonable times, to have access to and to copy and verify any records (A) bearing on whether the infant formula manufactured or held in the facility inspected meets the requirements of section 412, or (B) required to be maintained under section 412.

Sec. 704(b) Upon completion of any such inspection of a factory, warehouse, consulting laboratory, or other establishment, and prior to leaving the premises, the officer or employee making the inspection shall give to the owner, operator, or agent in charge a receipt describing the whole or in part of any filthy, putrid, or decomposed substance, or is otherwise unfit for food, a copy of the results of such analysis shall be furnished promptly to the owner, operator, or agent in charge.

Sec. 704(e) Every person required under section 519 or 520(g) to maintain records and every person who is in charge or custody of such records shall, upon request of an officer or employee designated by the Secretary, permit such officer or employee at all reasonable times to have access to and to copy and verify, such records.

Sec. 704(f)(1) An accredited person described in paragraph (3) shall maintain records documenting the training qualifications of the person and the employees of the person, the procedures used by the person for handling confidential information, the compensation arrangements made by the person, and the procedures used by the person to identify and avoid conflicts of interest. Upon the request of an officer or employee designated by the Secretary, the person shall permit the officer or employee, at all reasonable times, to have access to, to copy, and to verify, the records.

Sec. 512(j)(1) In the case of any new animal drug for which an approval of an application filed pursuant to subsection (b) is in effect, the applicant shall establish and maintain such records, and make such reports to the Secretary, of data relating to experience, including experience with uses authorized under subsection (a)(4)(A), and other data or information, received or otherwise obtained by such applicant with respect to such drug, or with respect to animal feeds bearing or containing such drug, as the Secretary may by general regulation, or by order with respect to such application, prescribe on the basis of a finding that such records and reports are necessary in order to enable the Secretary to determine, or facilitate a determination, whether there is or may be ground for invoking subsection (e) or subsection (m) of this section. Such regulation or order shall provide, where the Secretary decides it to be appropriate, for the examination, upon request, by the persons to whom such regulation or order is applicable, of similar information received or otherwise obtained by the Secretary.

(2) Every person required under this subsection to maintain records, and every person in charge or custody thereof, shall, upon request of an officer or employee designated by the Secretary, permit such officer or employee at all reasonable times to have access to and copy and verify such records.

Applicable sections of Parts F and G of Title III Public Health Service Act [42 U.S.C. 262-264] are quoted below:

Part F – Licensing – Biological Products and Clinical Laboratories

Sec. 351(c) "Any officer, agent, or employee of the Department of Health and Human Services, authorized by the Secretary for the purpose, may during all reasonable hours enter and inspect any establishment for the propagation or manufacture and preparation (Continued on Page 3)
of any virus, serum, toxin, antibiotic, vaccine, blood, blood component or derivative, allergenic product, or other product aforesaid for sale, barter, or exchange in the District of Columbia, or to be sent, carried, or brought from any State or possession into any other State or possession or into any foreign country, or from any foreign country into any State or possession."

Part F — * * * * * Control of Radiation.

Sec. 360 A (a) "If the Secretary finds for good cause that the methods, tests, or programs related to electronic product radiation safety in a particular factory, warehouse, or establishment in which electronic products are manufactured or held, may not be adequate or reliable, officers or employees duly designated by the Secretary, upon presenting appropriate credentials and a written notice to the owner, operator, or agent in charge, are thereafter authorized (1) to enter, at reasonable times any area in such factory, warehouse, or establishment in which the manufacturer's tests (or testing programs) required by section 358(h) are carried out and (2) to inspect, at reasonable times and within reasonable limits and in a reasonable manner, the facilities and procedures within such area which are related to electronic product radiation safety. Each such inspection shall be commenced and completed with reasonable promptness. In addition to other grounds upon which good cause may be found for purposes of this subsection, good cause will be considered to exist in any case where the manufacturer has introduced into commerce any electronic product which does not comply with an applicable standard prescribed under this subpart and with respect to which no exemption from the notification requirements has been granted by the Secretary under section 359(a)(2) or 359(e)."

(b) "Every manufacturer of electronic products shall establish and maintain such records (including testing records), make such reports, and provide such information, as the Secretary may reasonably require to enable him to determine whether such manufacturer has acted or is acting in compliance with this subpart and standards prescribed pursuant to this subpart and shall, upon request of an officer or employee duly designated by the Secretary, permit such officer or employee to inspect appropriate books, papers, records, and documents relevant to determining whether such manufacturer has acted or is acting in compliance with standards prescribed pursuant to section 359(a)."

(f) "The Secretary may by regulation (1) require dealers and distributors of electronic products, to which there are applicable standards prescribed under this subpart and the retail prices of which is not less than $50, to furnish manufacturers of such products such information as may be necessary to identify and locate, for purposes of section 359, the first purchasers of such products for purposes other than resale, and (2) require manufacturers to preserve such information Any regulation establishing a requirement pursuant to clause (1) of the preceding sentence shall (A) authorize such dealers and distributors to elect, in lieu of immediately furnishing such information to the manufacturer to hold and preserve such information until advised by the manufacturer or Secretary that such information is needed by the manufacturer for purposes of section 359, and (B) provide that the dealer or distributor shall, upon making such election, give prompt notice of such election (together with information identifying the notifier and the product) to the manufacturer and shall, when advised by the manufacturer or Secretary, of the need therefore for the purposes of Section 359, immediately furnish the manufacturer with the required information. If a dealer or distributor discontinues the dealing in or distribution of electronic products, he shall turn the information over to the manufacturer. Any manufacturer receiving information pursuant to this subsection concerning first purchasers of products for purposes other than resale shall treat it as confidential and may use it only if necessary for the purpose of notifying persons pursuant to section 359(a)."

Part G - Quarantine and Inspection

Sec. 360 B (a) It shall be unlawful—

(1) * * *

(2) * * *

(3) "for any person to fail or to refuse to establish or maintain records required by this subpart or to permit access by the Secretary or any of his duly authorized representatives to, or the copying of, such records, or to permit entry or inspection, as required or pursuant to section 360A."

Part G - Quarantine and Inspection

Sec. 361 (a) "The Surgeon General, with the approval of the Secretary, is authorized to make and enforce such regulations as in his judgment are necessary to prevent the introduction, transmission, or spread of communicable diseases from foreign countries into the States or possessions, or from one State or possession into any other State or possession. For purposes of carrying out and enforcing such regulations, the Surgeon General may provide for such inspection, fumigation, disinfection, sanitation, pest extermination, destruction of animals or articles found to be so infected or contaminated as to be sources of dangerous infection to human beings, and other measures, as in his judgment may be necessary."
<table>
<thead>
<tr>
<th><strong>DEPARTMENT OF HEALTH AND HUMAN SERVICES</strong></th>
<th><strong>FOOD AND DRUG ADMINISTRATION</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>1. DISTRICT ADDRESS AND PHONE NO.</td>
<td>6751 Steger Dr.</td>
</tr>
<tr>
<td></td>
<td>Cincinnati, OH 45237</td>
</tr>
<tr>
<td></td>
<td>(513)679-2700</td>
</tr>
</tbody>
</table>

| **2. NAME AND TITLE OF INDIVIDUAL** | **Michael A. Weston, Plant Manager** |
| **4. FIRM NAME** | **ABC Food Company** |
| **6. NUMBER AND STREET** | **3114 Mapleleaf Avenue** |
| **7. CITY AND STATE** | **Cincinnati, OH** |

| **3. DATE OF REQUEST** | **06/20/12** |
| **5. TIME OF REQUEST** | **8:30 AM** |
| **8. ZIP CODE** | **45213** |

Written demand for examination and/or copying of the records required by 21 CFR 113.100, 21 CFR 114 and 21 CFR 500.23 is hereby given, pursuant to 21 CFR 108.25(g), 21 CFR 108.35(h) and 21 CFR 500 for the records described below in order to verify the pH, adequacy of processing, the integrity of container closures, and the coding of the products processed by your firm.

**9. RECORDS NECESSARY**

All thermal process, production, and quality control / analytical records and maintenance records which may document any changes to the equipment or the thermal process mandated by 21 CFR 108, 113, and 114 [choose appropriate regulation, 113 LACF or 114 acidified] for all low acid canned foods and/or acidified food products [or specify product] which were produced by this firm since the last FDA inspection.

**10. SIGNATURE** (Food and Drug Administration Employee(s))

**Sidney H. Rogery**

**11. TITLE FDA EMPLOYEE**

Investigator
Written request is hereby given pursuant to 21 CFR 108.25(c)(3)(ii), 21 CFR 108.35(c)(3)(ii) and 21 CFR 500.23 for the information described below, concerning processes and procedures, which is deemed necessary by the Food and Drug Administration to determine the adequacy of the processes for products processed by your firm.

5. RECORDS NECESSARY

All documents and records mandated by 21 CFR 108 relating to or having a bearing on the adequacy of processes for all low acid canned foods and/or acidified food products [or specify product] that were produced in this firm since the last FDA inspection.
TO: William S. Gundstrom, Vice President, Production  

FIRM NAME: Topline Pharmaceuticals “T.L.P.”  

STREET ADDRESS: 2136 Elbe Place  

CITY, STATE AND ZIP CODE: Jackson, MN 55326  

TYPE OF ESTABLISHMENT INSPECTED: Tablet Repacker  

List your significant observations ranked in order of significance.  

See IOM 5.2.3, 5.2.3.1, 5.2.3.2, and 5.2.3.3
The observations of objectionable conditions and practices listed on the front of this form are reported:

1. Pursuant to Section 704(b) of the Federal Food, Drug and Cosmetic Act, or

2. To assist firms inspected in complying with the Acts and regulations enforced by the Food and Drug Administration.

Section 704(b) of the Federal Food, Drug, and Cosmetic Act (21 USC 374(b)) provides:

"Upon completion of any such inspection of a factory, warehouse, consulting laboratory, or other establishment, and prior to leaving the premises, the officer or employee making the inspection shall give to the owner, operator, or agent in charge a report in writing setting forth any conditions or practices observed by him which, in his judgement, indicate that any food, drug, device, or cosmetic in such establishment (1) consists in whole or in part of any filthy, putrid, or decomposed substance, or (2) has been prepared, packed, or held under insanitary conditions whereby it may have become contaminated with filth, or whereby it may have been rendered injurious to health. A copy of such report shall be sent promptly to the Secretary."
Screenshot - Resizing Pictures using Windows Explorer:

You can create resized copies of one or more selected pictures and store them in the current folder.

Select a size:
- Small (fits a 640 x 480 screen)
- Medium (fits a 800 x 600 screen)
- Large (fits a 1024 x 768 screen)
- Handheld PC (fits a 240 x 320 screen)

Advanced >>  OK  Cancel
Inserting a resized picture into Microsoft Word.
Screenshot - Using Microsoft Office Picture Manager to Resize a picture to 800 x 600 pixels.
EXHIBIT 5-9 INVESTIGATIONS OPERATIONS MANUAL 2016

Collect 12/100 tab bottles of lot DC-01234 as follow-up to violative EI of Pharma-Mix, Minneapolis, MN (FEI 3009001012) conducted on 9/31-10/05/2005. 30 cases were shipped to Drug Distributors Inc., 3910 Riverside St., Newark, NJ on 10/03/05 via Cross Country Express, Kansas City, MO. Invoice # 8328 10/05/05, B/L A-3026, 10-3-05.
### FORM FDA 482c (4/12)

#### Notice of Inspection

Notice of inspection is hereby given pursuant to Section 704(a)(1) of the Federal Food, Drug, and Cosmetic Act [21 U.S.C. 374(a)(1)]. Written request is hereby given to access and/or copy the records described below, pursuant to the Federal Food, Drug, and Cosmetic Act, Section 414(a) [21 U.S.C. 350c] and Title 21 Code of Federal Regulations, Section 1.361.

Applicable portions of Sections 704 and 414 of the Federal Food, Drug and Cosmetic Act (21 U.S.C. 374 and 350c) and Title 21 of the Code of Federal Regulations, are quoted below:

> "Sec. 704.(a)(1) For purposes of enforcement of this Act, officers or employees duly designated by the Secretary, upon presenting appropriate credentials and a written notice to the owner, operator, or agent in charge, are authorized (A) to enter, at reasonable times, any factory, warehouse, or establishment in which food, drugs, devices, or cosmetics are manufactured, processed, packed, or held, for introduction into interstate commerce; or after such introduction, or to enter any vehicle being used to transport or hold such food, drugs, devices, or cosmetics in interstate commerce; and (B) to inspect, at reasonable times and within reasonable limits and in a reasonable manner, such factory, warehouse, establishment, or vehicle and all pertinent equipment, finished and unfinished materials, containers and labeling therein. In the case of any person (excluding farms and restaurants) who manufactures, processes, packs, transports, distributes, holds, or imports foods, the inspection shall extend to all things therein (including records, packs, transports, distributes, holds, or imports such article shall, at the request of an officer or employee duly designated by the Secretary, permit such officer or employee, upon presentation of appropriate credentials and a written notice to such person, at reasonable times and within reasonable limits and in a reasonable manner, to have access to and copy all records relating to such article, and to any other article of food that the Secretary reasonably believes is likely to be affected in a similar manner, that are needed to assist the Secretary in determining whether the food is adulterated and presents a threat of serious adverse health consequences or death to humans or animals. (2) Use of or exposure to food of concern. --If the Secretary believes that there is a reasonable probability that the use of or exposure to an article of food, and any other article of food that the Secretary reasonably believes is likely to be affected in a similar manner, will cause serious adverse health consequences or death to humans or animals, each person (excluding farms and restaurants) who manufactures, processes, packs, distributes, receives, holds, or imports such article shall, at the request of an officer or employee duly designated by the Secretary, permit such officer or employee, upon presentation of appropriate credentials and a written notice to such person, at reasonable times and within reasonable limits and in a reasonable manner, to have access to and copy all records relating to such article, and to any other article of food that the Secretary reasonably believes is likely to be affected in a similar manner, that are needed to assist the Secretary in determining whether the food is adulterated and presents a threat of serious adverse health consequences or death to humans or animals. (3) Application.--The requirement under paragraphs (1) and (2) applies to all records relating to the manufacture, processing, packing, distribution, receipt, holding, or importation of such article maintained by or on behalf of such person in any format (including paper and electronic formats) and at any location."

> "Sec. 414(a) RECORDS INSPECTION. - (1) ADULTERATED FOOD. - If the Secretary has a reasonable belief that an article of food, and any other article of food that the Secretary reasonably believes is likely to be affected in a similar manner, is adulterated and presents a threat of serious adverse health consequences or death to humans or animals, each person (excluding farms and restaurants) who manufactures, processes, packs, distributes, receives, holds, or imports such article shall, at the request of an officer or employee duly designated by the Secretary, permit such officer or employee, upon presentation of appropriate credentials and a written notice to such person, at reasonable times and within reasonable limits and in a reasonable manner, to have access to and copy all records relating to such article, and to any other article of food that the Secretary reasonably believes is likely to be affected in a similar manner, that are needed to assist the Secretary in determining whether the food is adulterated and presents a threat of serious adverse health consequences or death to humans or animals. (2) Use of or exposure to food of concern. --If the Secretary believes that there is a reasonable probability that the use of or exposure to the food will cause serious adverse health consequences or death to humans or animals, each person (excluding farms and restaurants) who manufactures, processes, packs, distributes, receives, holds, or imports such article shall, at the request of an officer or employee duly designated by the Secretary, permit such officer or employee, upon presentation of appropriate credentials and a written notice to such person, at reasonable times and within reasonable limits and in a reasonable manner, to have access to and copy all records relating to such article, and to any other article of food that the Secretary reasonably believes is likely to be affected in a similar manner, that are needed to assist the Secretary in determining whether there is a reasonable probability that the use or exposure to the food will cause serious adverse health consequences or death to humans or animals. (3) Application.--The requirement under paragraphs (1) and (2) applies to all records relating to the manufacture, processing, packing, distribution, receipt, holding, or importation of such article maintained by or on behalf of such person in any format (including paper and electronic formats) and at any location."

> "321 C.F.R 1.361 What are the record availability requirements? When FDA has a reasonable belief that an article of food is adulterated and presents a threat of serious adverse health consequences or death to humans or animals, any records and other information accessible to FDA under section 414 or 704(a) of the act [21 U.S.C. 350c and 374(a)] must be made readily available for inspection and photocopying or other means of reproduction. Such records and other information must be made available as soon as possible, not to exceed 24 hours from the time of receipt of the official request, from an officer or employee duly designated by the Secretary of Health and Human Services who presents appropriate credentials and a written notice."

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**NOTICE OF INSPECTION - REQUEST FOR RECORDS**
1. Additive and batch weight known. Apply a straight edge to appropriate points on outside columns. Read ppm and/or percent additive where straight edge intersects central column.

2. Tolerance and batch weight known. Apply a straight edge to appropriate points on central and right-hand columns. Read the amount of additive in lbs. or gals where straight edge intersects the left-hand column.

For more precise determination of additives in the 1-500 ppm range, use Nomograph II.
### Food Additives Nomograph II

1. **Additive and Batch Weight Known.** Apply a straight edge to appropriate points on outside columns. Read ppm and/or percent additive where straight edge intersects central column.

2. **Tolerance and Batch Weight Known.** Apply a straight edge to appropriate points on central and right-hand columns. Read the amount of additive in lbs. or gals. where straight edge intersects the left-hand column.

<table>
<thead>
<tr>
<th>ADDITIVE (LBS OR GALS)</th>
<th>BATCH WEIGHT (LBS OR GALS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>1.0</td>
<td>8</td>
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<tr>
<td>0.9</td>
<td>7</td>
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<tr>
<td>0.8</td>
<td>6</td>
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<td>0.7</td>
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<td>8</td>
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<tr>
<td></td>
<td>5000</td>
</tr>
<tr>
<td></td>
<td>6000</td>
</tr>
</tbody>
</table>
### SUMMARY OF REGISTRATION AND LISTING REQUIREMENTS FOR THE MANUFACTURE OR DISTRIBUTION OF HUMAN PHARMACEUTICALS

<table>
<thead>
<tr>
<th>TYPE OF FIRM</th>
<th>REGISTRATION STATUS</th>
<th>LISTING STATUS</th>
<th>FACTS CODE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manufacturer [including homeopathic &amp; controlled drugs]</td>
<td>yes</td>
<td>yes</td>
<td>M</td>
</tr>
<tr>
<td>Contract Manufacturer</td>
<td>yes</td>
<td>yes</td>
<td>M</td>
</tr>
<tr>
<td>Own Label Distributor</td>
<td>no</td>
<td>yes</td>
<td>L</td>
</tr>
<tr>
<td>Wholesale Distributor</td>
<td>no</td>
<td>no</td>
<td>W-*</td>
</tr>
<tr>
<td>Own Label Repacker</td>
<td>yes</td>
<td>yes</td>
<td>R</td>
</tr>
<tr>
<td>Own Label Relabeler [including recirculizer]</td>
<td>yes</td>
<td>yes</td>
<td>Y</td>
</tr>
<tr>
<td>Contract Relabeler</td>
<td>yes</td>
<td>no</td>
<td>Y</td>
</tr>
<tr>
<td>Contract Testing Laboratory [dosage forms &amp; active ingredient release]</td>
<td>yes</td>
<td>no</td>
<td>C</td>
</tr>
<tr>
<td>Contract Testing Lab [doing non-release tests]</td>
<td>no</td>
<td>no</td>
<td>C</td>
</tr>
<tr>
<td>Contract Sub-Manufacturer</td>
<td>yes</td>
<td>no</td>
<td>M</td>
</tr>
<tr>
<td>IND Manufacturer [Clinical Drugs]</td>
<td>no</td>
<td>no</td>
<td>M</td>
</tr>
<tr>
<td>NDA and ANDA Manufacturer</td>
<td>yes</td>
<td>yes</td>
<td>M</td>
</tr>
<tr>
<td>Sponsor/Monitors/Clinical Investigator</td>
<td>no</td>
<td>no</td>
<td>4, 5, 6, 7</td>
</tr>
<tr>
<td>Contract Sterilizer</td>
<td>yes</td>
<td>no</td>
<td>0</td>
</tr>
<tr>
<td>Fulfillment Packager [adding substantive labeling]</td>
<td>yes</td>
<td>no</td>
<td>Y</td>
</tr>
<tr>
<td>Mail Order House [adding insubstantial labeling]</td>
<td>no</td>
<td>no</td>
<td>D</td>
</tr>
<tr>
<td>Printing House</td>
<td>no</td>
<td>no</td>
<td>None</td>
</tr>
<tr>
<td>Medical Gas Transfiller</td>
<td>yes</td>
<td>yes</td>
<td>MG</td>
</tr>
<tr>
<td>First Aid/Rescue Squad [transfilling for own use]</td>
<td>no</td>
<td>no</td>
<td>MG</td>
</tr>
<tr>
<td>Medical Gas Transfiller [operating out of a van]</td>
<td>yes</td>
<td>yes</td>
<td>MG</td>
</tr>
<tr>
<td>Contract Assembler</td>
<td>yes</td>
<td>no</td>
<td>M</td>
</tr>
<tr>
<td>Active Drug Substance Manufacturer</td>
<td>yes</td>
<td>yes</td>
<td>M</td>
</tr>
<tr>
<td>Excipient Drug Manufacturer</td>
<td>no</td>
<td>no</td>
<td>M</td>
</tr>
<tr>
<td>Manufacturer of Research Drugs</td>
<td>no</td>
<td>no</td>
<td>M</td>
</tr>
<tr>
<td>Drug Importer</td>
<td>no</td>
<td>no</td>
<td>A</td>
</tr>
<tr>
<td>Foreign Drug Manufacturer</td>
<td>yes</td>
<td>yes</td>
<td>M</td>
</tr>
<tr>
<td>Methadone Clinic</td>
<td>no</td>
<td>no</td>
<td>T</td>
</tr>
<tr>
<td>Retail Pharmacy</td>
<td>no</td>
<td>no</td>
<td>D</td>
</tr>
<tr>
<td>Salvage Operation</td>
<td>yes</td>
<td>no</td>
<td>X</td>
</tr>
<tr>
<td>Biopharmaceutical Clinical Facility</td>
<td>no</td>
<td>no</td>
<td>2</td>
</tr>
</tbody>
</table>

*Includes W, WA, WF, WR, and/or WZ
<table>
<thead>
<tr>
<th>Operation</th>
<th>Submit 510(k)</th>
<th>Register</th>
<th>List</th>
<th>COMPLY W/GMP</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Manufacture and distribute device</td>
<td>YES: 807.81(a)</td>
<td>YES 807.20</td>
<td>YES 807.20(a)</td>
<td>YES</td>
</tr>
<tr>
<td>2. Contract manufacturer who commercially distributes device for</td>
<td>NO: 807.81(a)</td>
<td>YES if domestic: 807.20(a)(2), YES if foreign 807.40(a)</td>
<td>YES 807.20(a)(2), YES if foreign 807.40(a)</td>
<td>YES</td>
</tr>
<tr>
<td>specifications developer</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3a. Contract manufacturer who meets the definition of finished</td>
<td>NO</td>
<td>YES 807.20(c)(1)</td>
<td>YES 807.20(c)(1)</td>
<td>YES</td>
</tr>
<tr>
<td>device manufacturer per 21 CFR 820.3(i)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3b. Contract manufacturer who does not meet the definition of</td>
<td>NO</td>
<td>YES 807.20(c)(1)</td>
<td>YES 807.20(c)(1)</td>
<td>NO</td>
</tr>
<tr>
<td>finished device manufacturer per 21 CFR 820.3(i) (e.g., component</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>manufacturer, subassembler)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Manufacturer modifies device or new intended use and distribute</td>
<td>NO: preamble no. 17 &amp; 18 FR 820.23/77, YES: 807.81(a)(3) with significant change in device or use</td>
<td>YES 807.20(a)</td>
<td>YES 807.20(a)</td>
<td>YES</td>
</tr>
<tr>
<td>(domestic distributor)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Distribute U.S. Made device: no specification initiation</td>
<td>NO: 807.85(b)</td>
<td>NO: 510(g)(4) of act, 807.20(c)(3)</td>
<td>NO</td>
<td>NO</td>
</tr>
<tr>
<td>(domestic distributor)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Specification initiator and distribute only</td>
<td>YES: 807.81(a)</td>
<td>YES 807.20(a)(1) preamble no. 5, FR 8-23-77</td>
<td>YES 807.20(a)(1)</td>
<td>YES: 820.181, etc.</td>
</tr>
<tr>
<td>7. Specification consultant only; no distribution</td>
<td>NO</td>
<td>NO: preamble no. 5, FR 8-3-77</td>
<td>NO</td>
<td>NO</td>
</tr>
<tr>
<td>8. Relabeler or repacker: distribute under own name</td>
<td>NO: 807.85(b); no change to device or existing labeling</td>
<td>YES 807.20(a)(3)</td>
<td>YES 807.20(a)(3) preamble no. 7, FR 8-25-78</td>
<td>YES</td>
</tr>
<tr>
<td>9. Kit assembler using prelabeled &amp; prepackaged devices only</td>
<td>NO: no change in device or existing labeling</td>
<td>YES 807.20(a)</td>
<td>YES 807.20(a)</td>
<td>NO</td>
</tr>
<tr>
<td>10. Kit assembler changes intended use (801.4) of prepackaged/prelabeled</td>
<td>YES: 807.81(a)</td>
<td>YES 807.20(a)(2)</td>
<td>YES 807.20(a)(2)</td>
<td>YES: 820.120, 820.130, etc.</td>
</tr>
<tr>
<td>devices only</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. Kit assembler changes prepackaged/prelabeled devices</td>
<td>NO: if no significant change to labeling or device: otherwise</td>
<td>YES 807.81(a)(3)(i)</td>
<td>YES 807.20(a)(3)</td>
<td>YES</td>
</tr>
<tr>
<td>12. Manuf. Accessory, component and package &amp; label for health purpose to</td>
<td>YES: 807.81(a)</td>
<td>YES 807.20(a)(5) preamble no. 77, FR 8-25-78</td>
<td>YES 807.20(a)(5)</td>
<td>YES</td>
</tr>
<tr>
<td>end user</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>13. Manuf. Components &amp; dist. Only to finished device mfr.</td>
<td>NO: 807.81(a)</td>
<td>NO: 807.85(a)</td>
<td>NO</td>
<td>Use as guide: 820.1</td>
</tr>
<tr>
<td>14. Contract mfr. Of subassembly or component (see no. 12, accessory)</td>
<td>NO</td>
<td>NO</td>
<td>NO</td>
<td>Primary mfr. must see that GMP is met preamble no. 33, FR 7-21-78</td>
</tr>
<tr>
<td>15. Contract packager or labeler</td>
<td>NO</td>
<td>NO</td>
<td>NO</td>
<td>Primary mfr. must see that GMP is met 21 CFR 820.50</td>
</tr>
<tr>
<td>16. Contract Sterilizer</td>
<td>NO</td>
<td>YES if domestic 807.20(a)(2), YES if foreign 807.40(a)</td>
<td>YES 807.20(a)(2), YES if foreign 807.40(a)</td>
<td>YES</td>
</tr>
<tr>
<td>17. Manufacture custom device (domestic or foreign)</td>
<td>NO: 807.85(a)(1)(A)(2)</td>
<td>YES 807.20(a)(2)</td>
<td>YES 807.20(a)(2)</td>
<td>YES: also see 520(b); 520(f)</td>
</tr>
<tr>
<td>18. U.S. Establishment who manufactures for export only</td>
<td>NO</td>
<td>YES 807.20(a)(2)</td>
<td>YES 807.20(a)(2)</td>
<td>YES</td>
</tr>
<tr>
<td>19. Foreign manufacturers and all foreign establishments</td>
<td>YES: 807.81 foreign mfr. has primary responsibility, but may delegate to an init. Dist.</td>
<td>YES 807.40(a)</td>
<td>YES 807.40(a)</td>
<td>YES</td>
</tr>
<tr>
<td>20. Initial distributor/importer of device</td>
<td>YES: 807.81(a) or 807.85(b) unless 510(k) has been filed by foreign manufacturer or another init. Dist</td>
<td>YES 807.20(a)(4)</td>
<td>NO: enforcement discretion used for 807.22(c)</td>
<td>YES 807.3(d), 820.198, 820.100, 820.200, etc.</td>
</tr>
<tr>
<td>21. Installer-mfr.'s agent</td>
<td>NO</td>
<td>NO</td>
<td>NO</td>
<td>YES 820.170</td>
</tr>
<tr>
<td>22. Installer-user</td>
<td>NO</td>
<td>NO</td>
<td>NO</td>
<td>NO: for x-ray see 1020.30(d) report</td>
</tr>
<tr>
<td>23. Device being investigated under ide</td>
<td>Exempt: 812.1(a)</td>
<td>NO</td>
<td>NO: 807.40(c)</td>
<td>Exempt per 812.1(a), except for Design Control per 820.30</td>
</tr>
<tr>
<td>24. Mfr. Buys manufacturing rights for device (see no. 4)</td>
<td>NO: preamble 18 FR 8-23-77 only if same type of manuf. equip. is used and no significant change to device</td>
<td>YES 807.20(a)(2) if not already registered</td>
<td>Send letter to FDA per 807.30(b)(5) &amp; 807.25</td>
<td>YES</td>
</tr>
<tr>
<td>25. Reprocessor of single use device</td>
<td>YES</td>
<td>YES 807.20</td>
<td>YES 807.20</td>
<td>YES</td>
</tr>
<tr>
<td>26. Foreign exporter of device (device manufactured in foreign country)</td>
<td>YES: (original manufacturer's 510(k) maybe used)</td>
<td>YES 807.40(a)</td>
<td>YES 807.40(a)</td>
<td>YES 820.1(a)(2) YES</td>
</tr>
</tbody>
</table>
PROFILING A FIRM’S CGMP/QS COMPLIANCE STATUS

Table 5-14.1 Quick Reference Guide

<table>
<thead>
<tr>
<th>Review Status</th>
<th>Profile Status</th>
<th>Data Entry Role</th>
<th>Remarks Field</th>
<th>Remarks Status Field</th>
<th>Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial</td>
<td>Further Action Indicated</td>
<td>IB</td>
<td>Review and date Ex: “Referred to CB mm/dd/yy”</td>
<td></td>
<td>EI is potentially OAI</td>
</tr>
<tr>
<td></td>
<td>Acceptable</td>
<td>IB</td>
<td>Usually no Remarks required.</td>
<td></td>
<td>EI is NAI or VAI.</td>
</tr>
<tr>
<td>In Review</td>
<td>Pending</td>
<td>CB</td>
<td>Recommended enforcement or alternative action; with date as well as review and date. Ex: “Recommend WL: Under review by [CB/Center]”</td>
<td></td>
<td>Enforcement or alternative action recommended.</td>
</tr>
<tr>
<td>Final</td>
<td>Other</td>
<td>IB/CB</td>
<td>Enter the action firm is operating under Ex: “Consent Decree (CD) for CGMP (Current Good Manufacturing Practices)/QS (Quality Systems) violations signed on mm/dd/yy.” If the CD includes a sunset clause/date, add to Remarks. or “AIP invoked on mm/dd/yy.” When the firm is operating under CD/Injunction/AIP (Application Integrity Policy) and the CGMP/QS EI is: NAI or VAI, then “Acceptable (AC)” or the inspection is OAI and further enforcement action* is taken, then the Remarks Status is “Unacceptable (UN).”</td>
<td></td>
<td>Firm is operating under a CD or AIP, and a subsequent CGMP EI has occurred. Enforcement Action* may involve medically necessary products or be process or product specific. In this case, such conditions should be reflected in Remarks field (see 3.10 &amp; 3.11(2)).</td>
</tr>
<tr>
<td></td>
<td>Acceptable</td>
<td>IB/CB</td>
<td>No outstanding OAI inspections No compliance actions</td>
<td></td>
<td>NAI and VAI inspections; or OAI inspections where no enforcement action was taken and/or was downgraded to VAI.</td>
</tr>
</tbody>
</table>
Unacceptable CB Outstanding OAI Inspections

*NOTE: A final profile status of unacceptable must be supported with regulatory action as recognized in the Field Management Directive sec 86 (FMD 86). This would include a Warning Letter, seizure, injunction, or prosecution based on CGMP deficiencies. A regulatory meeting or an untitled letter for CGMP deficiencies are not considered enforcement actions for purposes of an unacceptable profile status.

Table 5-14.2 Example of a Maintain Profiles screen

<table>
<thead>
<tr>
<th>Firm</th>
<th>FEI: 1234567890</th>
<th>Name: Cloudy Day, Inc.</th>
<th>Address: 1234 Sunshine Lane</th>
<th>City: Rockslide</th>
<th>State: WA</th>
<th>Zip Code: 24567</th>
</tr>
</thead>
<tbody>
<tr>
<td>Profile Classes</td>
<td>Profile Description</td>
<td>Operation Type</td>
<td>Last GMP Date</td>
<td>Last Other Insp Date</td>
<td>Last Final Status</td>
<td>Pending Action Ind</td>
</tr>
<tr>
<td>COS</td>
<td>Computer Software</td>
<td>Manufacturer</td>
<td>11/14/2009</td>
<td>Others</td>
<td></td>
<td>Add</td>
</tr>
<tr>
<td>ELE</td>
<td>Electronic assembly</td>
<td>Manufacturer</td>
<td>11/14/2009</td>
<td>Others</td>
<td></td>
<td>Delete</td>
</tr>
<tr>
<td>PRF</td>
<td>Plastic or rubber fabrication</td>
<td>Manufacturer</td>
<td>11/14/2009</td>
<td>Others</td>
<td></td>
<td>Discontinue</td>
</tr>
</tbody>
</table>

Current Profile Status

Date Type: GMP Inspection Date: 10/05/2009

Initial: In Review: Final:

Further Action

Pending

Others

Remarks

Investigator A

Compliance Officer B

Compliance Officer B

Recommended By

SEA-IB-DA

SEA-CB

SEA-CB

District Code

Record Initial

Record In Review

Record Final

Delete Status

Example of a firm operating under a Consent Decree where the current CGMP inspection (10/5/009) is OAI.

5-14.1 Introduction

Firm profiles provide a snapshot of the firm’s compliance status with CGMP or QS regulations. Profile status is monitored for domestic and foreign firms that manufacture, repack, label/relabel, sterilize, or test drug, medical device, or biological products.

5-14.2 Purpose

Firm profiles provide the compliance status as well as an inventory of product categories covered during a CGMP/QS inspection and are used to support:

- The Government Wide Quality Assurance Program (GWQAP).
- External users such as state and local regulatory authorities and foreign government agencies.
- Other FDA operations such as drug product approvals, export certificates and imports.

5-14.3 Instructions

5-14.3.1 Pre-Inspection Preparation

To obtain a comprehensive history of the firm you are going to inspect go to ORADSS Domestic Reports folder named Establishment History Report and select EHR02_Firm Info and run the report entering the FEI and start and end dates for the period you want reported. Make sure that a final status has been entered for all PCs for the previous inspection. If you find that one or more PCs have an initial status but not a final status, bring this to the attention of your supervisor and finalize prior entering any updates.

5-14.3.2 Firm’s Operations

For profile purposes, the firm’s operation type can be either as a single entity or in combination with other operations. Look at all the possibilities in the drop down menu before making a selection. Some selections allow for multiple operations. See below for examples:

a. Specification Developer Only versus Specification Developer Also

When a firm is a specification developer and they do not manufacture any medical products onsite, select profile class code, SPD, and the Operation Type, “Specification Developer Only.”

When a firm is a specification developer and they do onsite manufacturing of medical products which are not the subject of the specifications developed, select SPD with the Operation Type “Specification Developer Also” and select the appropriate profile class (PC) of the products they manufacture with Operation Type “Manufacturer.”

b. Control Testing Lab Only versus Control Testing Lab Also

When a firm is a contract control testing laboratory only and does not manufacture medical products, select profile class code, CTL, and Operation Type, “Control Testing Lab Only.”
When a firm is a (contract) control testing laboratory and manufactures its own medical products onsite, select the appropriate profile class code(s) that defines its operation, e.g., CTX for drugs, CTD for devices, or CTB for biologics (or a combination) and Operation Type, “Control Testing Laboratory Also.” Select also the PCs that define the class of products they manufacture and Operation Type, “Manufacturer.”

c. Veterinary Drugs Also versus Veterinary Drugs Only

When a firm manufactures both veterinary and human drugs, select the appropriate profile class code(s) that defines its operation then select the Operation Type Veterinary Drug Also. When a firm manufactures veterinary drugs only, select the appropriate profile class code(s) that define its operation then select the Operation Type Veterinary Drugs Only.

For other changes in operations, or discontinuing profile required operations of FDA regulated products, update the establishment type industry code information on the MARCS/FACTS Firm Management Services (FMS) screen.

5-14.3.3 Maintain Profiles Screen

When entering profile information, it is important to access the Maintain Profiles screen properly as accessing a profile screen incorrectly will result in data quality errors.

The correct way for Field Offices and Centers to access the Profile screen is to access MARCS (Mission Accomplishments and Regulatory Compliance Services) FACTS (Field Assignments and Compliance Tracking System) database from Inside.FDA’s ORA Applications, FACTS web link – clicking on Go takes you to the FDA MARCS Application Production screen. From the menu toolbar, choose Navigate, scroll down to Investigative Operations, move over and click on Inspections. Enter the FEI and click ExecQry to bring up the Maintain Inspection Results screen. From the toolbar, click the profile icon or click Options and scroll down to Profiles. You are now ready to enter/update the Maintain Profile record.

If there is a need to search for a firm through the FACTS Firms Detail screen, select the Inspection button. This will take you to the Maintain Inspection Results screen and from there use Options to reach Profiles. Do not enter profile information via this screen.

NOTE: Information entered/updated through this screen will not be linked to an inspection. Entering and/or updating profile information from this screen is the cause of many profile data errors and problems, such as duplications and/or non-finalized profiles which cause problems with future updates of inspectional information.

5-14.3.4 Previous Inspection Profile

It is important that the profile for the previous inspection be complete with a final profile status for each PC before updating the profile for the current inspection. If this is not done, a banner will appear saying “Initial data already exists,” and it will not be possible to close the current inspection in FACTS on the Maintain Inspection Results screen.

5-14.3.5 Firm Information

The Maintain Firm, Maintain Inspection Results, and Maintain Profiles screens should agree in firm name, address, and FEI number. If not, or if the firm has a name or address change, the change must be made on the Maintain Firm screen. For questions, contact your District FACTS Profile Monitor. For foreign firms, contact your trip planner concerning discrepancies. See IOM Directory, ORA Field Program Monitors for contact information.

5-14.3.6 Inspection Coverage of Profile Class Codes

When a CGMP/QS systems based inspection is performed, coverage should reflect the overall state of control for the firm’s operations. For this reason, the PCs should reflect all product classes produced by the firm and covered during the inspection, even if they are not directly covered. For example, if a firm is a drug manufacturer and a CGMP/QS systems based inspection is performed, then all PCs should be updated for all products produced by the firm.

When a firm manufactures more than one commodity, e.g., drugs and devices, and the inspection covers only the drug systems, then only update the PCs that represent the drug commodity. See 5-14.7 for more information about profile classes and codes.

5-14.3.7 Discontinue and Delete Buttons

Proper use of the Discontinue and the Delete buttons: Discontinue button – The PCs should be discontinued if a firm goes out-of-business or no longer manufactures a drug, device, or biologic product. Delete button - PCs and data entered in error can and should be deleted prior to clicking the save button and exiting the screen.

NOTE: If you save incorrect data before realizing it and you cannot delete it, contact the GWQAP Team for assistance. See 5-14.4 for Contact Information.

5-14.3.8 CGMP Inspection and Other Toggle Buttons

The CGMP Inspection toggle button is automatically activated when the Profile Required field is checked on the Maintain Inspection Results screen. The Other radio button should not be used for profiling purposes.
5-14.3.9 Initial, In Review, and Final
As reflected in Table 5-14.1 above, profile status should be entered as follows:

**Initial**: Normally entered by the Investigator. Potentially OAI inspections should be immediately entered as FI and NAI/VAI as AC.

**In Review**: Pending should be entered by the Compliance Officer as soon as the record is received for review.

**Final**: AC should be entered by the Supervisor for NAI/VAI inspections; UN should be entered by the Compliance Officer for OAI inspections when a regulatory action has been taken.

NOTE: The Status Date automatically records the date that the information is entered or updated in Initial, In Review, and Final Profile Status. It is important to maintain the integrity of the profile information by not changing this date.

**Foreign firms**: The districts enter the initial status only and the appropriate center enters the final profile class.

5-14.3.10 Final Profile Status
It is important for the Field and Centers to understand that final profile status should be promptly entered when a final agency decision has been made. Profiles should not be held in Pending status if the District or Center decides that the course of action is to not take enforcement action as defined by FMD-86, and, instead, re-inspect.

NOTE: This represents a change in procedure. Previously, an In Review “Pending” status was permitted until a re-inspection was made. Now, unless the District or Center plans to accelerate the re-inspection on an elevated priority basis to reach a final decision, the District or Center should close out the profile status. A final status of UN must be supported with a regulatory action. See 5-14.3.10.3. Please contact GWQAP Team with any concerns. For contact information, see 5-14.4.

5-14.3.10.1 Other Status
Other should be entered as the final profile status for all profile class codes when a firm is operating under a CD or AIP. See Tables 5-14.1 & 5-14.2 above for more information.

5-14.3.10.2 Acceptable Status
AC should be entered as the final profile status when an inspection is classified as NAI or VAI and the firm is not operating under a CD or AIP. See Table 5-14.1 above for more information. If an OAI is not supported by an enforcement action, it is entered as AC as defined in Field Management Directive (FMD)-86.

5-14.3.10.3 Unacceptable Status
UN should be entered as the final profile status when there is an outstanding OAI inspection. Refer to FMD-86 for final classification instructions.

5-14.3.10.3.1 Continuation of Unacceptable Status
A UN status along with the regulatory action taken may be carried forward from one inspection to the next when the follow-up inspection reveals the firm had not addressed the violations identified in the original OAI inspection or an enforcement action. In this case, it is important the Remarks field note this condition. See 5-14.3.11 Remarks field for more information.

5-14.3.10.3.2 Changing from Unacceptable to Acceptable Status
A UN status may be changed to AC when the agency’s review of the firm’s response to a warning letter reveals the firm’s corrective actions adequately address the violations identified, a re-inspection for verification may or may not be warranted. The Remarks field must note the reason for the change.

5-14.3.11 Remarks Status Field
The Remarks Status field is used mainly to indicate the compliance status of a current inspection while the firm operates under a CD or AIP. See Tables 5-14.1 & 5-14.2 for more information and examples.

It may also be used to indicate an exception to the general compliance status. The profile status when under a CD will be “Others.” The Remarks Status Field will show the current compliance inspection status (AC/UN). The Remarks Field will note that the firm is operating under a CD (include date and any information required concerning the current inspection.

5-14.3.12 Remarks Field
The Remarks field is a narrative field to be used as often as needed to:

1. Track the status of any potential or completed enforcement or alternative action with dates. This may include an explanation for a continuation of an UN final profile status from one inspection to the next when the follow up inspection reveals the firm’s corrective actions were found inadequate. See Table 5-14.1 above or 5-14.4 below for more information and accessing the ORA/OEIO/DCS intranet site, respectively.

2. Indicate when a firm is operating under a CD or AIP with date. Note when there are specific conditions such as product(s) subject to the CD or AIP. This information must remain in Remarks for each PC until the CD/AIP is vacated or revoked.

3. Identify product(s) covered when using the catch all PCs MIS for devices, BMI for biologics and NEC for drugs; and

4. Indicate where a sterilization process(es) takes place such as onsite at the manufacturer, or offsite by a contract sterilizer. If offsite, include the name, address, and FEI of the contract sterilizer.

NOTE: After entering the information once, a copy and paste method can be used to update the Remarks field for each profile class involved as follows:
a. Highlight the narrative text by clicking in the Remarks field.
b. Select CTRL C to copy.
c. Select CTRL V to paste.

5-14.3.13 Change in Operational Status
When the MARCS/FACTS “operational status” of a profiled firm changes, an assessment must be made to determine if the firm is still required to be profiled. If, for example, the firm goes out of business the profile and registration fields must be discontinued and cancelled respectively prior to setting the operational status to Out-of-Business (OOB). Remember to uncheck the Profiled Required box from the List of Values (LOV) under Operational Status and select Out-of-Business (OOB) or Not Official Establishment Inventory (NOE). For profile changes other than OOB or not OEI firms refer to IOM Exhibit 5-14.3.7. Also select the Maintain Profiles screen and discontinue each profile class as follows:

1. If appropriate, verify that the registration is cancelled by the registration monitor.
3. Navigate to the “Enter Additional Firm Details” screen in Firm Management Services (FMS).
4. Click the Edit link under Status.
5. Select the appropriate operational status from the LOV. For example, select OOB for an out of business firm.
6. Verify the Workload Obligation is appropriately set. For example, for OOB it should be set to N.
7. For OOB and NOE firms, uncheck Profile Required.
8. Select the Profile tab at the top of the “Firm Details” screen.
9. In MARCS/FACTS, select update button in the “Maintain Profile” screen and edit each PC.
10. Save changes and close window.

5-14.3.14 Firm Merge
Before attempting to merge two or more firm records, always check to ensure all profile class codes have been finalized. Do not attempt to merge if the profile status is left in Initial or In Review. Merging firms where the profile classes are not finalized will cause problems that can only be resolved through FDA’s information technology service department and consumes Agency resources. For questions, contact the GWQAP Team. See 5-14.4 below for contact information.

5-14.3.15 Troubleshooting
Troubleshooting information may be found at DCIQA’s intranet site. See 5-14.4 for intranet site location.

5-14.4 Contact Information
Go to ORA/OEIO/Division of Compliance Systems and select the appropriate Program Area.

NOTE: The COMSTAT Team has been changed to the GWQAP Team. To contact DCS, GWQAP Team

Email: GWQAP@FDA.HHS.GOV

5-14.5 Data Quality Assurance Projects
Our GWQAP stakeholders, including the Department of Veterans Affairs (VA), the Department of Defense (DoD) [through the Defense Supply Center Philadelphia (DSCP)], as well as several Local, State, and Foreign Governments, use an external view of FACTS profiles to help them make procurement decisions for medical products. Since these stakeholders can view only the latest acceptable or unacceptable final profile status, profile classes must be finalized.

Use the active down arrow at the top of the Maintain Inspections Results screen to view the previous inspection and profiles covered. Make sure a final status has been entered for all PCs for the previous inspection. If you find one or more PC has an initial status but not a final status, bring this to the attention of your supervisor.

5-14.6 Establishment Profile Criteria
Table 5-14.6.1 Device, Biologic, Drug, and Veterinary Establishments TO Profile

<table>
<thead>
<tr>
<th>Manufacturer</th>
<th>Makes a new or a changed product from one or more ingredients.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remanufacturer</td>
<td>Processes, conditions, renovates, repackages, restores, or performs any other act to a finished device that significantly changes the device’s performance or safety specifications or intended use.</td>
</tr>
<tr>
<td>Reprocessor</td>
<td>Performs remanufacturing operations on a single use device.</td>
</tr>
<tr>
<td>Packer/Repacker</td>
<td>Packs a product or products into different containers without making any changes in the form of the product.</td>
</tr>
<tr>
<td>Labeler/Relabeler</td>
<td>An establishment which affixes the original labeling to a product or changes in any way the labeling on a product without affecting the product or its container.</td>
</tr>
<tr>
<td>Contract Sterilizers</td>
<td>Performs sterilization or irradiation of products or components of products regulated by FDA on a contract basis.</td>
</tr>
<tr>
<td>Control Testing Laboratories</td>
<td>Performs production quality control work related to products regulated by FDA on a contract basis.</td>
</tr>
<tr>
<td>Assemblers of Medical Device Kits</td>
<td>Responsible for assembling finished devices into medical device kits.</td>
</tr>
<tr>
<td>Tissue Establishments</td>
<td>Only tissue establishments inspected as device firms under the Quality</td>
</tr>
</tbody>
</table>
System regulations.

| Specification Developer | Initiates or develops specifications for a device that is distributed under the establishment's own name but is manufactured by a second person. |

Table 5-14.6.2 Establishment and Operations NOT to Profile

| Blood Banks |
| Methadone Clinics |
| Manufacturers of "Research Use Only" Products |
| Pharmacies (including pharmacy compounders) and Retail firms |
| Distributors |
| Plasmapheresis Centers |
| Custom Device Manufacturers |
| Veterinary Medical Device Firms |
| X-ray Assemblers |
| Mammography Clinics |
| Manufacturers of General Purpose Articles (Devices) |
| Physicians’ Offices, Hospitals and Clinics |
| Laser Light Shows/Television and Microwave Oven Manufacturers |
| Sun tanning Establishments |
| Device Component Manufacturers |
| Clinical Investigators/Bioresearch Monitoring |
| Tissue firms inspected under Good Tissue Practices |
| Any Non-GMP Inspection |

5-14.6.3 Pre-Approval Inspections

Pre-Approval inspections that cover the firm's systems (Quality Control + 1) should be treated like any other CGMP or QSIT inspection and should be profiled. When a pre-approval inspection is the initial inspection of firm and results in the firm not being approved to market within the U.S., the firm should not be profiled. The investigator should uncheck the profile required box on the Maintain Inspection Results screen.

NOTE: If the decision not to find the firm acceptable is reversed by the Center review, the Center is responsible for contacting the District to have the firm profiled. The District will inform the Center when the update is complete so the Center can then enter the Final classification.

When a pre-approval inspection finds problems affecting the product approval, but they do not affect the overall CGMP/QSR compliance status of the firm, the Profile Status should be entered as acceptable.

Information regarding withholding the approval of the product should be annotated on the Maintain Inspection Results screen in the Remarks field under District Decisions.

5-14.7 Profile Classes and Codes

The profile system is based upon product categories or classes, and is not product specific. Select the most appropriate profile class(es) to describe the product(s) the firm manufactures or otherwise processes.

When describing devices, often more than one class is needed to describe the operations/assembly involved in the device. A rule of thumb is to think of the composition of the device and then select the profile classes that define the make-up of that device and its assembly. For example a catheter and needle unit is profiled as MTL (metal fabrication and assembly) and PRF (plastic or rubber fabrication and assembly). A Cutter, orthopedic cast, 110 volt AC-DC, is profiled as MTL, PRF and ELE (electrical) For devices that have software and are operated by computer, codes COS (software) and COH (computer hardware) should be added.

SPD (specification developer) should be used if a firm only develops the design and specifications and has the device manufactured by someone else. In this case use only SPD, do not include other profile classes unless the firm also manufactures other medical products on-site.

Catch-all codes: MIS for devices, NEC and CRU for drugs, and BMI for biologics can be used when product does not fit into any product class identified by the list of PCs. When using these codes, identify the type of product in the Remarks field for that code. If the product is a sterile product, don’t forget to include the appropriate sterilization PC and identify if onsite or offsite.

When a product or products have been transferred from one Center to another, discontinue the profile class from the former Center representing that product if that profile class is no longer needed for any remaining products, and add a profile class which represents the new Center.

NOTE: Some Drug definitions have been updated.

5-14.7.1 Profile Class Codes

For more information, contact your District Profile Monitor or DCIQA. See 5-14.4 for contact information.

Table 5-14.7.1.1 Biologics

<table>
<thead>
<tr>
<th>BIOLOGICS</th>
<th>Definitions</th>
</tr>
</thead>
<tbody>
<tr>
<td>AEV</td>
<td>ANTITOXINS AND ANTIVENINS</td>
</tr>
<tr>
<td>AFP</td>
<td>ANIMAL DERIVED FRACTIONATION PRODUCTS</td>
</tr>
<tr>
<td>ALP</td>
<td>ALLERGENIC PRODUCTS</td>
</tr>
<tr>
<td>BGR</td>
<td>BLOOD GROUPING REAGENTS</td>
</tr>
<tr>
<td>BMI</td>
<td>BIOLOGICAL PRODUCTS NOT ELSEWHERE CLASSIFIED (Blood collection bags with anti-coagulant, plasma volume expanders, Limulus Amebocyte Lysate (LAL) test kit, etc.; Note specific products(s) in Remarks field)</td>
</tr>
<tr>
<td>CBS</td>
<td>COMPUTER BIOLOGICAL SOFTWARE</td>
</tr>
<tr>
<td>CGT</td>
<td>CELL AND GENE THERAPY PRODUCTS</td>
</tr>
<tr>
<td>CTB</td>
<td>CONTROL TESTING LABORATORY “ALSO”</td>
</tr>
<tr>
<td>HFP</td>
<td>HUMAN DERIVED FRACTIONATION PRODUCTS</td>
</tr>
<tr>
<td>RBD</td>
<td>RECOMBINANT ANALOGUES OF BLOOD DERIVATIVE PRODUCTS</td>
</tr>
</tbody>
</table>
## Table 5-14.7.1.2 Devices

<table>
<thead>
<tr>
<th>Code</th>
<th>Class</th>
<th>Profile</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADM</td>
<td>TRAJECTORY</td>
<td>ADM</td>
<td>AEROSOL DISPENSED MEDICATION</td>
</tr>
<tr>
<td>CBI</td>
<td>RECOMBINANT</td>
<td>API</td>
<td>RECOMBINANT/Non-RECOMBINANT PROTEIN DS OF BIOLOGIC ORIGIN</td>
</tr>
<tr>
<td>CEX</td>
<td>STARTING</td>
<td>INTERMEDIATE</td>
<td>Derived from plant/animal EXTRACTION</td>
</tr>
<tr>
<td>CFN</td>
<td>NON-STERILE</td>
<td>API</td>
<td>Fermentation</td>
</tr>
<tr>
<td>CFS</td>
<td>STERILE</td>
<td>API</td>
<td>By FERMENTATION</td>
</tr>
<tr>
<td>CHG</td>
<td>CAPSULES</td>
<td>PROMPT RELEASE</td>
<td></td>
</tr>
<tr>
<td>CRU</td>
<td>NON-STERILE</td>
<td>STARTING/INTERMEDIATE/NEC (not Plant/Animal)</td>
<td></td>
</tr>
<tr>
<td>CRX</td>
<td>STERILE</td>
<td>STARTING/INTERMEDIATE/NEC (not Plant/Animal)</td>
<td></td>
</tr>
<tr>
<td>CSG</td>
<td>CAPSULES</td>
<td>SOFT GELATIN</td>
<td></td>
</tr>
<tr>
<td>CSN</td>
<td>NON-STERILE</td>
<td>API</td>
<td>By CHEMICAL SYNTHESIS</td>
</tr>
<tr>
<td>CSS</td>
<td>STERILE</td>
<td>API</td>
<td>By CHEMICAL SYNTHESIS</td>
</tr>
<tr>
<td>CTR</td>
<td>CAPSULES</td>
<td>MODIFIED RELEASE</td>
<td></td>
</tr>
<tr>
<td>CTX</td>
<td>CONTROL</td>
<td>TESTING LABORATORIES ALSO - Drugs</td>
<td></td>
</tr>
<tr>
<td>CXA</td>
<td>PURIFIED API</td>
<td>DERIVED FROM PLANT/ANIMAL EXTRACTION</td>
<td></td>
</tr>
<tr>
<td>EXC</td>
<td>EXCIPIENT</td>
<td>(also referred to as inactive ingredient)</td>
<td></td>
</tr>
<tr>
<td>GAS</td>
<td>MEDICAL</td>
<td>GAS</td>
<td>(includes liquid oxygen)</td>
</tr>
<tr>
<td>LIQ</td>
<td>NON-STERILE</td>
<td>LIQUID</td>
<td>(other than suspensions &amp; emulsions)</td>
</tr>
<tr>
<td>LVP</td>
<td>LARGE VOLUME</td>
<td>PARENTERALS</td>
<td></td>
</tr>
<tr>
<td>NEC</td>
<td>NOT ELSEWHERE</td>
<td>CLASSIFIED FINISHED DRUG</td>
<td></td>
</tr>
<tr>
<td>OIN</td>
<td>OINTMENTS</td>
<td>NON-STERILE</td>
<td>(including creams, jelly, paste, etc.)</td>
</tr>
<tr>
<td>PET</td>
<td>POSITRON</td>
<td>EMISSION</td>
<td>TOMOGRAPHY</td>
</tr>
<tr>
<td>POW</td>
<td>NON-STERILE</td>
<td>POWDERS</td>
<td>(includes oral and topical)</td>
</tr>
<tr>
<td>SES</td>
<td>SUSPENSIONS</td>
<td>AND EMULSIONS</td>
<td>(NON PARENTERALS)</td>
</tr>
<tr>
<td>SLQ</td>
<td>STERILE</td>
<td>LIQUID</td>
<td>(other than suspensions &amp; emulsions)</td>
</tr>
<tr>
<td>SON</td>
<td>STERILE</td>
<td>OINTMENT</td>
<td></td>
</tr>
<tr>
<td>SPW</td>
<td>STERILE</td>
<td>POWDER</td>
<td></td>
</tr>
<tr>
<td>SUP</td>
<td>SUPPOSITORIES</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SVL</td>
<td>SMALL VOLUME</td>
<td>PARENTERALS</td>
<td>(Lyophilized)</td>
</tr>
<tr>
<td>SVS</td>
<td>STERILE</td>
<td>FILLED SMALL VOLUME PARENTERAL DRUGS</td>
<td></td>
</tr>
<tr>
<td>SVT</td>
<td>TERMINALLY</td>
<td>STERILIZED</td>
<td>SMALL VOLUME PARENTERALS</td>
</tr>
<tr>
<td>TCD</td>
<td>TERMINALLY</td>
<td>STERILIZED</td>
<td>SMALL VOLUME PARENTERALS</td>
</tr>
<tr>
<td>TCT</td>
<td>TABLETS</td>
<td>PROMPT RELEASE</td>
<td></td>
</tr>
<tr>
<td>TDP</td>
<td>TRANSDERMAL</td>
<td>PATCHES</td>
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## Table 5-14.7.3 Drugs and Veterinary

<table>
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<tr>
<th>Code</th>
<th>Class</th>
<th>Profile</th>
<th>Definition</th>
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<tr>
<td>ADM</td>
<td>TRAJECTORY</td>
<td>ADM</td>
<td>AEROSOL DISPENSED MEDICATION</td>
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## Table 5-14.7.4 Miscellaneous

<table>
<thead>
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<th>Profile</th>
<th>Definition</th>
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<td>ADM</td>
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## Table 5-14.7.5 Special Veterinary
### 5-14.8 Abbreviations and Definitions

<table>
<thead>
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<th>Abbreviation</th>
<th>Definition</th>
<th>Reference</th>
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<tr>
<td>AC</td>
<td>Acceptable</td>
<td>Table 5-14.1 Quick Reference Guide</td>
</tr>
<tr>
<td>AIP</td>
<td>Application Integrity Policy</td>
<td>Table 5-14.1</td>
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<tr>
<td>CB</td>
<td>Compliance Branch</td>
<td>Table 5-14.1</td>
</tr>
<tr>
<td>CD</td>
<td>Consent Decree</td>
<td>Table 5-14.1</td>
</tr>
<tr>
<td>CGMP</td>
<td>Current Good Manufacturing Practice</td>
<td>Table 5-14.1</td>
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<tr>
<td>CTRL</td>
<td>Control</td>
<td>5-14.3.11</td>
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<tr>
<td>DCS</td>
<td>Division of Compliance Systems</td>
<td>5-14.4</td>
</tr>
<tr>
<td>DoD</td>
<td>Department of Defense</td>
<td>5-14.5</td>
</tr>
<tr>
<td>EI</td>
<td>Establishment Inspection</td>
<td>Table 5-14.1</td>
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<tr>
<td>FACTS</td>
<td>Field Assignments and Compliance Tracking System</td>
<td>5-14.3.2</td>
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<td>FDA Establishment Identifier</td>
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<td>FI</td>
<td>Further Action Indicated</td>
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<td>FMD</td>
<td>Field Management Directive</td>
<td>5-14.3.9</td>
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<td>GWQAP</td>
<td>Government-Wide Quality Assurance Program</td>
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<tr>
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<tr>
<td>OOB</td>
<td>Out of Business</td>
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### Compliance Achievements

**Firm Reported By**
- **Firm:**
- **Organization:** CHI-DO
- **Address:**
- **Employee:**
- **Home District:** CHI-DO

**Corrective Actions**

<table>
<thead>
<tr>
<th>Product Code</th>
<th>PAC</th>
<th>Problem Type</th>
<th>Corrective Action</th>
<th>Verification Date</th>
<th>Reporting Organization</th>
<th>Correcting Organization</th>
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**Reason for Correction**

**Remarks**

**Description**

**Linked Operations**

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<th>Op ID</th>
<th>Type of Operation</th>
<th>Unique ID</th>
<th>Accomplishing Organization</th>
<th>Performing Organization</th>
<th>Status</th>
<th>Status Date</th>
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</thead>
<tbody>
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</table>
### Inspection Results

#### Name:

- **Compliance Status:**
- **Completion Date:**

#### District Decisions

- **Final Decision:**
  - **Decision Date:**
  - **Decision Type:**

#### Products Covered

<table>
<thead>
<tr>
<th>Product Code</th>
<th>Establishment Type</th>
<th>Product Description</th>
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</table>

### IB Suggested Actions

- **Warning Letter**

### Referrals

<table>
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<tr>
<th>Org Name</th>
<th>Mail Code</th>
<th>Referral Reason</th>
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### Refusals

- **Inspection Refusal:**
  - **Refusal to permit photography**
  - **Refusal to permit collection of samples**

### Samples Collected

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<tr>
<th>Sample Number</th>
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### Recall Numbers

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### Related Complaints

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### Maintain Products Covered

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<th>Description</th>
<th>Last Covered Date</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>F E I: 30300000702</td>
<td>Name: Standard Seafood Co.</td>
<td>Milwaukee, WI 53204</td>
<td>United States</td>
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<td></td>
<td></td>
<td>Address: 3050 Telegraph Rd.</td>
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<tr>
<td></td>
<td></td>
<td>Manufacturer</td>
<td>Shrimp &amp; Prawns, Breaded, Paper, Packaged</td>
<td>10/17/2005</td>
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<tr>
<td></td>
<td>16 J G</td>
<td>Manufacturer</td>
<td>Shrimp &amp; Prawns, Plastic, Synth, Heat Treated</td>
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<tr>
<td></td>
<td>16 E H T</td>
<td>Manufacturer</td>
<td>Clams, Nonflex Plastic, Packaged Food (Not Commercial)</td>
<td></td>
</tr>
</tbody>
</table>
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Date: (Enter Date)

To: Supervisory Investigator

From: Jane Smith
Investigator, Florida District

Subject: Special Investigation

Text of Investigation

ENDORSEMENT

TO:

Signature of Supervisory Investigator (or designee)

O: Home District Responsible Firm
cc: State or Federal Agency
cc: Accomplishing District
cc: Program Monitor (if applicable)
cc: Consumer Complaint Coordinator (if applicable)
Screenshot showing location of Reimbursable check box:
Format of EIR Addendum

Establishment Inspection Report
XYZ Corporation
Chesterton, IN 46304

FEI: 1234567890
EI Start: 08/30/2010
EI End: 09/24/2010

ADDENDUM

Clearly define what section(s) and pages are being revised; what information is being added or clarified.

Sidney H. Rogers, Investigator
DET-DO
South Bend RP