U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES
U.S. Food and Drug Administration

FDA Emergency Operations Plan
Version 3.0

July 2019

OFFICE OF EMERGENCY MANAGEMENT
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A MESSAGE FROM THE ACTING COMMISSIONER TO CENTER/OFFICE LEADERSHIP

FDA’s mission to protect and promote the public health requires that FDA remain prepared to deal with a wide variety of natural and manmade threats that involve, impact, or require the use of FDA-regulated products. FDA has significant experience responding to such threats, and our country has benefited greatly from the support of our dedicated staff during such emergencies.

In order to ensure that we remain prepared to respond effectively to threats, FDA’s Emergency Operations Plan (EOP) needs to be accurate and current. The EOP summarizes how the agency coordinates information and resources during emergencies.

We are issuing the first major update to the EOP since 2014. The updated EOP reflects organizational changes within the agency, particularly the Office of Regulatory Affairs program alignment and the addition of FDA’s Executive Committee in the response process. These process changes were assessed and validated by external independent study conducted in the Fall of 2018.

While each threat is unique, the magnitude of recent events — such as the increased number of hurricanes — has required FDA to engage in new ways to ensure product safety and availability for our fellow citizens. Recent emergency responses have demonstrated the critical nature of coordinating resources across FDA and prioritizing our efforts. We have been able expand our response capacity by leveraging staff expertise from across the agency, and have been able to better identify the right approaches and solutions by collaborating with Federal partners, industry, and the community.

Key to our EOP is timely communication and, in that regard, the revised plan facilitates the rapid dissemination of information, especially to affected consumers, about our response and the steps toward recovery. Our ongoing training programs ensure that we are able to work effectively as a unified team to mitigate, respond to, and recover from disasters.

Catastrophic events, when they occur, challenge FDA to respond effectively to such crises and provide relief to affected Americans. The work of the FDA staff who put in countless hours in support of these efforts — on top of their other mission-critical work at the agency — is deeply appreciated. Along with the agency’s senior leaders, I am fully committed to FDA’s role in support of the nation’s efforts to recover from any disaster.

Any questions regarding this EOP can be directed to the Office of Emergency Management/Emergency Planning, Exercises, and Evaluation Staff (OCMExerciseandEvaluations@fda.hhs.gov).

Ned Sharpless, M.D.
Acting Commissioner of Food and Drugs
An electronic version of the FDA EOP and its annexes can be found on the Inside FDA website at
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U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES
U.S. Food and Drug Administration

FDA Emergency Operations Plan
Version 3.0

July 2019

OFFICE OF EMERGENCY MANAGEMENT
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A. INTRODUCTION

For purposes of this plan and its supporting annexes:

An incident is defined as: An occurrence or event, natural or manmade, that requires the attention of the U.S. Food and Drug Administration (FDA). At the incident level, FDA is aware of the circumstances, is monitoring and assessing the situation, and is determining whether it has regulatory authority over the incident. FDA may initiate response operations as described in this plan as appropriate. Any incident may evolve into an emergency.

An emergency is defined as: An unforeseen occurrence or a combination of circumstances that poses a significant risk to public health and may include, but is not limited to, the safety, efficacy, and security of human and veterinary medicines, biological products, medical devices, our Nation’s food supply, cosmetics, products that emit radiation, and tobacco products that call for immediate actions by FDA staff.

Since each emergency is unpredictable and dynamic, and any incident has the potential to escalate into an emergency, this plan will use the terms “incident” and “emergency” interchangeably.

Emergencies and disasters, whether natural or manmade, accidental or intentional, have the potential to cause adverse health and safety effects for large segments of the human and animal populations. In order to mitigate the consequences of such incidents, FDA must possess the resources and capabilities necessary to prevent, prepare for, protect against, and rapidly and effectively respond to and recover from all hazards. A planned and coordinated approach to emergency operations by FDA organizational components in support of Federal, State, local, tribal, and territorial (SLTT) government, with assistance when appropriate from foreign counterparts and international partners, can save lives and ensure that critical public health and medical needs are met.

This FDA Emergency Operations Plan (EOP) and its annexes are an all-discipline, all-hazards plan that establishes a single, comprehensive framework for FDA’s management of incidents. It provides the measures, operating structures, roles and responsibilities, and mechanisms for direction and coordination of FDA resources before, during, and after disease outbreaks, terrorist attacks and other criminal acts, natural disasters, and any other incidents associated with FDA-regulated products that pose a risk to human or animal health.

This FDA EOP is compatible with the scalable, flexible, and adaptable Federal government emergency coordinating structures of the National Response Framework (NRF); is consistent with the concepts, principles, and terminology of the National Incident Management System (NIMS);¹ and fulfills the requirements of:

- National Preparedness Goal (NPG)
- National Preparedness System
- National Planning Frameworks
- Federal Interagency Operational Plans

¹ For more information on the NRF and NIMS, refer to the “Authorities and References” section of this EOP.


This FDA EOP is to be used to guide FDA in conducting response operations for all types of incidents.

### A.1 MISSION

FDA is responsible for protecting the public health by ensuring the safety, efficacy, and security of human and veterinary medicines, biological products, medical devices, our Nation’s food supply, cosmetics, and products that emit radiation and by regulating tobacco products. Additionally, FDA advances public health by helping to speed innovations that make foods safer and medical products more effective, safer, and more affordable and by helping the public get the accurate scientific information it needs to use medical products and foods to improve their health.

FDA must continue to meet its mission during emergency and disaster situations, under adverse conditions, and/or from alternate locations as needed. In addition to day-to-day activities, FDA activities may expand to include performance of specific incident-related functions, such as:

- Assisting and supporting the U.S. Department of Health and Human Services (HHS) in public health- and medical-related efforts to prevent, respond, mitigate, and recover from an incident.
- Conducting inspections and investigations and assessing damage to FDA-regulated industry in impacted areas.
- Coordinating the tracing (forward and backward) of the distribution of any potentially contaminated FDA-regulated products and initiating seizures or recalls as appropriate.
- Collecting and analyzing product samples.
- Providing technical assistance or subject matter expertise related to food and feed, human and animal drugs, medical devices, biologics, radiation-emitting devices, and tobacco products.
- Conducting assessments of food retail establishments in impacted areas.
- Conducting assessments of tobacco product retail establishments nationwide.
- Conducting reviews of all adverse event reports related to FDA-regulated products.
- Assisting with surveillance efforts to determine product integrity of foods, pharmaceuticals, medical supplies, equipment, and tobacco products.
- Authorizing the use of an unapproved medical product (drug, biologic, and device) or the unapproved use of an approved medical product (e.g., emergency use authorization [EUA]).
- Issuing safety alerts, health information advisories, warnings, and advice and guidance to consumers and industry.
- Requiring product manufacturers to make safety-related changes to prescribing information or labeling.
- Detaining or removing contaminated or unfit merchandise from the market and/or restricting those marketing it.
A.2 **Purpose**

The purpose of the FDA EOP is to provide for a coordinated and consistent agency approach to preparing for, preventing, protecting against, mitigating, responding to, and recovering from incidents involving or impacting FDA-regulated products.

To accomplish this, the FDA EOP:

- Serves as the single, overarching FDA-wide operational plan to address the full spectrum of natural and technological hazards and terrorist threats and the “umbrella plan” into which all supporting agency emergency plans, procedural documents, and other guidance integrate.
- Defines the FDA emergency operating structure and assigns essential tasks to all FDA organizational components involved in prevention, protection, mitigation, response, and recovery efforts.
- Provides mechanisms for vertical and horizontal command, control, coordination, and communications.
- Integrates FDA emergency response operations into the Federal coordinating structure and ensures consistency with nationally recognized incident management policy and guidance.

A.3 **Scope and Applicability**

The FDA EOP covers the full range of complex and constantly changing requirements in anticipation of or in response to all “incidents” that FDA manages or participates in, including the following:

- Complaints, adverse events, recalls, or unintentional contamination involving FDA-regulated products that present a threat of serious adverse health consequences or death to humans or animals.
- Natural disasters (e.g., hurricanes, tornadoes, severe storms, floods, fires, earthquakes, volcanic eruptions, tsunamis, landslides).
- Naturally occurring disease and foodborne illness outbreaks, epidemics, and pandemics.
- Manmade accidents, such as hazardous materials releases or spills; air, land, or water contamination; and utility outages.
- Terrorist or criminal acts, including the threat or intentional use of chemical, biological, radiological, nuclear, or high-yield explosive (CBRNE) weapons against human or animal populations or FDA-regulated products.

The FDA EOP establishes intra- and interagency mechanisms for FDA involvement in domestic and international incident management operations. These mechanisms include coordinating structures and processes for incidents requiring agency support for consumer protection and to other Federal agencies; States and territories, tribal nations, and local governments; foreign governments; and international organizations. It is applicable to all FDA headquarters and field organizational components that may be required to provide assistance or conduct emergency operations in the context of actual or potential incidents. In these cases, FDA may use the Incident Command System (ICS) to facilitate command and coordination.

A.4 **Planning Assumptions**

The FDA EOP is based on the following planning assumptions:

- Incidents, including large-scale emergencies and major disasters, will require full coordination of FDA operations and resources, and may:
− Occur at any time with little or no warning in the context of a general or specific threat or hazard.
− Involve one or more FDA organizational component(s) and span a single or multiple districts.
− Require significant information sharing, resource coordination, and/or assistance across FDA organizational components, Federal and SLTT agencies, the private sector, and foreign governments.
− Result in numerous casualties, fatalities, and displaced people; property loss; significant damage to the environment; and disruption of economy and normal life-support systems, essential public services, and critical infrastructures.
− Require extremely short-notice FDA response times and prolonged, sustained recovery operations and support activities.

- All FDA emergency-related activities will be initiated and conducted in accordance with the principles, concepts, and terminology established within NIMS.
- Regardless of incident characteristics or requirements, FDA continues to be responsible for consumer products under its jurisdiction while coordinating response operations with other interagency public health and medical partners.
- An emergency or disaster that overwhels the capabilities of State and local governments may require FDA and/or other Federal agencies to assist in meeting public health and safety needs. FDA assets will supplement the response, as directed or requested, when an incident affects an FDA-regulated product or requires the provision of subject matter expertise or use of specific medical countermeasures. As such, FDA will participate in an Incident Command or Unified Command (UC) system to manage multi-agency emergencies.
- Contamination of the Nation’s human and animal food or medical products supply may initially be indistinguishable from a naturally occurring event. Moreover, depending upon the particular agent and associated signs or symptoms, several days or weeks could pass before authorities suspect terrorism may be the cause.
- Response to a chemical, biological, radiological, or nuclear (CBRN) incident suspected of being deliberate in origin or a terrorist act requires consideration of special law enforcement and homeland security requirements as well as international legal obligations and requirements.
- The combined expertise and capabilities of government at all levels, industry, and nongovernmental organizations (NGOs) will be required to respond to incidents of catastrophic proportions. During such periods, FDA will provide emergency support as directed by the Commissioner, the Secretary of HHS, and/or the Secretary of Homeland Security through the coordinating mechanisms established within the NRF and National Disaster Recovery Framework (NDRF).
- Supporting documentation (e.g., standard operating procedures [SOPs], operations and procedural manuals, field guidance) has been developed and made available to designated FDA emergency staff to provide detailed instructions during performance of specific functions or incident-related actions.
- As part of its Emergency Support Function (ESF) #8 functions, FDA can provide SLTT agencies with activities related to regulated product support. For example, FDA can provide assistance to State and local agencies in ensuring the safety of food products at retail food establishments and drug and medical products in pharmacies.
To ensure the capability to implement the FDA EOP, each FDA headquarters and field organizational component tasked with emergency roles and responsibilities, as identified in this EOP basic plan or any of its annexes, may develop and maintain individual emergency plans and procedures that identify the critical and time-sensitive missions, functions, assignments, and processes to be performed during all hazards. These documents shall be consistent with this EOP and be available to all FDA personnel.

A.5 Activation

While many parts of this EOP may be used routinely to manage FDA’s emergency operations, the full plan will be activated when certain events occur such as, but not limited to, the following:

- A determination has been made by the Commissioner of Food and Drugs (the Commissioner) or other FDA senior official(s) as delegated by the Commissioner that FDA emergency actions are warranted to prevent, prepare for, protect against, mitigate, respond to, or recover from a threat or hazard.
- The Secretary of HHS (the Secretary) has directed FDA to provide emergency support to maintain the safety, efficacy, and security of drugs, biologics, medical devices, radiation-emitting devices, the food supply, or cosmetics.
- The Secretary has directed FDA to provide emergency support to protect the public health in the event of tobacco product adulteration.
- The Secretary of Homeland Security has raised the National Terrorism Advisory System (NTAS) alert level.
- The President has declared that an emergency or a major disaster exists within an affected State or States as defined in the Robert T. Stafford Disaster Relief and Emergency Assistance Act (Stafford Act), P.L. 100-707, as amended, and that NRF coordinating mechanisms, in whole or in part, have been activated.
- The President has declared a national emergency exists under the National Emergencies Act, 50 United States Code (U.S.C.) 1601-1651, in the event that the Nation is threatened by crisis, exigency, or emergency circumstances (other than natural disasters, war, or near-war situations).
- The Secretary of HHS may, under Section 319 of the Public Health Service Act (PHSA), determine that (a) a disease or disorder presents a public health emergency; or (b) that a public health emergency, including significant outbreaks of infectious disease or bioterrorist attacks, otherwise exists.

A.5.1 Activation of EOP Annexes

A decision on whether an EOP annex is to be implemented in response to an incident is made by the Office of Emergency Management (OEM) Director and senior officials of the Center(s) involved, or in a meeting requested by the Commissioner, a Center Director, the Associate Commissioner for Regulatory Affairs (ACRA), the Director of OEM, Associate Commissioner for Counterterrorism, or some other top official specifically to discuss a particular incident or problem. If an annex(es) is to be activated, the OEM Director will notify the involved FDA parties (through electronic mail, if available).

\[2\] District Directors and Center/Office Directors may activate their individual organizational component’s EOP without activation of the FDA EOP.
A.6 SUPERSEDERENCE

This FDA EOP (version 3.0) supersedes the FDA EOP (version 2.0) and its annexes, dated March 2014.
B. CONCEPT OF OPERATIONS

The scope of FDA emergency operations is as wide as the array of products the agency regulates. FDA actions impact the Nation’s quality of life on a daily basis through its regulation of food, drugs, medical devices, radiation-emitting products, biologics, veterinary drugs, and tobacco products. For this reason, FDA’s successful and efficient handling of emergencies and disasters is especially important in fulfilling its mission to maintain not only the public’s health, but also its trust and confidence. The following concept of operations (CONOPS) describes the principal authorities governing agency emergency functions and the phases within which FDA conducts incident-related operations.3

B.1 EMERGENCY AUTHORITIES

Statutory authorities relevant to FDA emergency preparedness and response are described below under the general categories of safety and medical countermeasures. In addition, certain FDA regulations in Title 21 of the Code of Federal Regulations (CFR) may be particularly applicable in an emergency.

- Safety of FDA-regulated products:
  - Under PHSA, 42 U.S.C. 262, FDA will only approve a biologics license application upon a demonstration that the biological product is safe, pure, and potent. In addition, under PHSA, 42 U.S.C. 264, FDA may issue and enforce regulations necessary to prevent the spread of communicable disease.

- Medical, including specific authorities related to medical countermeasures:
  - Under the FD&C Act, FDA generally approves drugs and devices that have been shown to be safe and effective. Under PHSA, FDA licenses biological products based upon a determination that they are safe, pure, and potent.
  - Section 561 of the FD&C Act (Expanded Access to Unapproved Therapies and Diagnostics) allows FDA to permit the treatment of a patient with an investigational medical countermeasure under certain circumstances, even though the safety and effectiveness of the drug have not been fully established.
  - Section 564 of the FD&C Act, as established by the Project BioShield Act of 2004 and amended by the Pandemic and All-Hazards Preparedness Act of 2013, the 21st Century Cures Act of 2016, and P.L. 115-92, allows FDA to authorize the emergency use of unapproved drug(s), device(s), or biological product(s) for certain types of emergencies if there is a declaration justifying the need for the product and FDA determines the criteria for issuance have been met.
  - Section 564A of the FD&C Act allows FDA to take certain actions to authorize certain uses of FDA-approved medical products that otherwise could render the product unapproved or misbranded, such as provide Emergency Dispensing orders to allow dispensing without individual prescriptions, extensions of expiration dates, and waivers of CGMPs.

3 For detailed information on the actions performed during specific incidents, refer to the FDA EOP Incident Annexes.
B.2 EMERGENCY OPERATIONS PHASES

The following phases comprise the entire spectrum of FDA emergency operations: Prevention, Protection, Mitigation, Response, and Recovery. Although emergency operations may involve each of these phases over the course of any multitude of incidents, the nature and severity of an event and the FDA organizational component(s) responding will determine the specific order, actions, and responsible parties required for each.4

B.2.1 Prevention, Protection, and Mitigation

FDA closely integrates the prevention, protection, and mitigation phases. These three phases are mission areas under the NPG and defined as follows:

- **Prevention** includes those capabilities necessary to avoid, prevent, or stop a threatened or actual act of terrorism. It is focused on ensuring we are optimally prepared to prevent an imminent attack within the United States.

- **Protection** includes capabilities to safeguard the homeland against acts of terrorism and manmade or natural disasters. It is focused on actions to protect the citizens, residents, visitors, and critical assets, systems, and networks against the greatest risks to our Nation in a manner that allows our interests, aspirations, and way of life to thrive. We will create conditions for a safer, more secure, and more resilient Nation by enhancing protection through cooperation and collaboration with all sectors of society.

- **Mitigation** includes those capabilities necessary to reduce loss of life and property by lessening the impact of disasters. It is focused on the premise that individuals, the private sector, communities, critical infrastructure, and the Nation as a whole are made more resilient when the consequences and impacts, the duration, and the financial and human costs to respond and recover from adverse incidents are all reduced.

The “prevention, protection, and mitigation” phase of FDA emergency operations includes those actions taken to avoid an incident or intervene to stop an incident from occurring, or to mitigate its effects on FDA-regulated products and the community, during periods of increased risk or heightened threat conditions.5 It involves immediate steps to protect consumers; ensure the safety and defense of food and animal feed, medical products (e.g., drugs, vaccines, blood donations, and diagnostic devices), cosmetics, field operations, and toxicological research; and apply intelligence and other information to a range of directed countermeasures. Upon preliminary determination of an incident, FDA conducts the following preventive/protective/mitigation measures:

- Conducting scientific *vulnerability assessments* of different categories of food to determine the most serious risks of intentional contamination.

- Providing *preventive training tools* to government, industry, and other stakeholders.

- Increasing *surveillance and detection* of adverse events, disease outbreaks, natural disasters, and other emerging public health concerns.

- Heightening *consumer product protection* measures against the adverse effects of naturally occurring and manmade hazards.

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4 Refer to *Section C*, “Organization and Assignment of Responsibilities,” of this EOP for an overview of FDA organizational component roles and responsibilities during all hazards. Refer to the FDA EOP Incident Annexes for incident-specific Center/Office actions.

5 Examples of high-risk situations include designated National Special Security Events (NSSEs), developing public health emergencies and natural disasters, and NTAS alert levels.
• Facilitating the availability of medical countermeasures that prevent, diagnose, or treat diseases or conditions caused by CBRN agents, as well as medical products in response to natural disasters.

• Increasing surveillance of FDA-regulated products used to prevent or mitigate adverse health effects related to the actual or potential incidents.

FDA works closely with DHS and other Federal agencies to coordinate the overall national effort to enhance the protection of critical infrastructure and key resources of the Nation, including food and agriculture defense. DHS serves as the coordinator of the Food and Agriculture Sector with the Government Coordination Council (GCC). The sector is a public-private partnership that combines expertise from several Federal agencies as well as SLTT officials (representing agriculture, public health, and veterinary services) and the private sector (more than 100 trade associations and individual firms).

B.2.1.1 Surveillance, Detection, and Alert

Within the context of this plan, increased risk-based “surveillance and detection” is defined as an increase in the frequency, quantity, or detail of the ongoing systematic collection, analysis, and interpretation of public health data essential to the planning, execution, and evaluation of FDA emergency operations, closely integrated with the timely dissemination of these data to those responsible for prevention and control. FDA uses accumulated data from a variety of sources (e.g., international, Federal, and SLTT public health agencies; consumer complaints; regulatory inspections, investigations, and sampling; laboratory testing) for passive and active surveillance of regulated products and other public health concerns during day-to-day situations and for targeted preventive actions. The focus of these systems is to detect a “signal” to allow for additional information gathering and analysis or to track and trace a suspected or confirmed event within the agency’s jurisdiction. They support FDA’s primary mission, including identifying and reviewing adverse events and tracking and tracing product problems, and are used to facilitate counterterrorism and product safety and security activities as appropriate.

The Office of Emergency Operations (OEO) is in a constant state of readiness during routine agency operations, maintaining 24 hours a day, 7 days a week (24/7) monitoring capability for surveillance and detection. OEM/OEO staff members assist with the detection of signals either as the direct point of contact (POC) for outside stakeholders or in support of the Centers and Office of Regulatory Affairs (ORA) and participate in meetings to identify emerging threats. OEM’s geographic information system (GIS) is another tool utilized to analyze health information and FDA activity data for spatially occurring patterns on a local, regional, or national level. FDA’s After-Hours Emergency Call Center serves as an additional resource during non-duty hours, monitoring complaints and reports of problem products and triaging phone calls. Pre-identified topics are assessed by OEM/OEO staff to determine potential public health threats.

In addition, several databases and electronic information exchanges enable responsible FDA organizational components to collect and report out-of-the-ordinary information, monitor potential and ongoing situations, and determine whether agency emergency response activities are warranted (Table B-1).
### Table B-1. Surveillance and Detection Systems Utilized by FDA

<table>
<thead>
<tr>
<th>System</th>
<th>Description</th>
<th>Sponsoring Agency</th>
<th>FDA POC</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDA’s Adverse Event Reporting System (FAERS)</td>
<td>The FAERS is a computerized information database designed to support FDA’s post-marketing safety surveillance program for all approved drug and therapeutic biological products. FDA uses FAERS to monitor for new adverse events and medication errors that might occur with these marketed products.</td>
<td>FDA</td>
<td>CDER</td>
</tr>
<tr>
<td>Biological Product Deviation Reporting (BDPR) System</td>
<td>The BDPR System reports errors and accidents in manufacturing of products, including testing, processing, packing, labeling, or storage, or with the holding or distribution of a licensed biological product or a blood or a blood component, in which the safety, purity, or potency of a distributed product may be affected. Also, it includes deviations in manufacturing of human cells, tissues, and cellular and tissue-based products that relate to the prevention of communicable disease transmission, contamination, or other unexpected events.</td>
<td>FDA</td>
<td>CBER</td>
</tr>
<tr>
<td>BioWatch</td>
<td>BioWatch is a detection system for the release of biological agents in the air through a comprehensive protocol of monitoring and laboratory analysis.</td>
<td>CDC</td>
<td>OEM</td>
</tr>
<tr>
<td>CDRH Product Availability (Shortages) Database</td>
<td>The Shortages Database identifies and monitors supplies of certain devices that are or have the potential to be in demand.</td>
<td>FDA</td>
<td>CDRH</td>
</tr>
<tr>
<td>CFSAN Adverse Event Reporting System (CAERS)</td>
<td>The CAERS receives and monitors all post-marketing surveillance adverse events reports that directly affect CFSAN. These adverse event reports include foods, dietary supplements, cosmetic products, and food and color additives.</td>
<td>FDA</td>
<td>CFSAN</td>
</tr>
<tr>
<td>Counterfeit Alert Network (CAN)</td>
<td>The CAN is a coalition of health professional and consumer groups to disseminate educational guidance and alert messages about counterfeit drug incidents and measures to take to prevent exposure.</td>
<td>FDA</td>
<td>ORA/OCI</td>
</tr>
<tr>
<td>Electronic Laboratory Exchange Network (eLEXNET)</td>
<td>The eLEXNET is an integrated, web-based information network that allows health officials at multiple government agencies engaged in food safety activities to coordinate and share laboratory analysis findings. It provides the necessary infrastructure for an early warning system that identifies potentially hazardous foods and enables health officials to assess risks and analyze trends. eLEXNET is the data capture and communication system for the Food Emergency Response Network (FERN) and is supported by the USDA and DoD.</td>
<td>FDA</td>
<td>ORA/OO</td>
</tr>
<tr>
<td>Emergency Operations Network – Incident Management System (EON-IMS)</td>
<td>The EON-IMS serves as the central hub for exchanging and relaying all incident-related information within the agency. Managed by OEM, this system integrates multiple data streams from other electronic systems (such as the FERN, eLEXNET, and Epidemic Information Exchange [Epi-X]) and from FDA laboratories and investigators and external agencies, into a coherent fashion during critical decision points. The EON-IMS creates a safety net that significantly reduces the probability that incidents will prevent FDA from accomplishing its objectives and minimizes the impact of these events on normal operations.</td>
<td>FDA</td>
<td>OEM</td>
</tr>
<tr>
<td>Epidemic Information Exchange (Epi-X)</td>
<td>The Epi-X is a secure, web-based communications network that provides public health officials with up-to-the-minute information, reports, alerts, and discussions about terrorist events, toxic exposures, disease outbreaks, and other public health events. It provides a flexible search interface for researching outbreaks and unusual health events and for tracking information for outbreak investigations and response.</td>
<td>CDC</td>
<td>OEM</td>
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</tbody>
</table>

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6 For more information on EON-IMS, refer to Section E, “Communications and Information Management,” in this EOP.
<table>
<thead>
<tr>
<th>System</th>
<th>Description</th>
<th>Sponsoring Agency</th>
<th>FDA POC</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Foodborne and Diarrheal Diseases Outbreak Summaries</strong></td>
<td>FDA receives the weekly summaries on outbreaks from the CDC Foodborne and Diarrheal Diseases Branch to document ongoing investigations, current outbreaks, State(s) where the outbreak is occurring, and the serotype of the causal microorganism.</td>
<td>CDC</td>
<td>CFSAN</td>
</tr>
<tr>
<td><strong>Foodborne Diseases Active Surveillance Network (FoodNet)</strong></td>
<td>The FoodNet is a collaborative project of the CDC, 10 State Emerging Infections Program sites, the FSIS of the USDA, and FDA. It consists of active surveillance for foodborne diseases and related epidemiologic studies designed to help public health officials better understand the epidemiology of foodborne diseases in the United States.</td>
<td>CDC, USDA, FDA, State Health Departments</td>
<td>CFSAN</td>
</tr>
<tr>
<td><strong>Food Emergency Response Network (FERN)</strong></td>
<td>FERN is a network of Federal and State laboratories that analyze food samples in the event of a biological, chemical, nuclear, or radiological incident or large-scale contamination. FERN Federal partners include FDA, USDA, CDC, DHS/U.S. Customs and Border Protection (CBP), and the EPA.</td>
<td>FDA and USDA</td>
<td>ORA</td>
</tr>
<tr>
<td><strong>Health Alert Network (HAN)</strong></td>
<td>The HAN provides rapid and timely access to emergent health information and evidence-based practices and procedures for effective public health preparedness, response, and service on a 24/7 basis. It includes all 50 States, three large city health departments, three county health departments, eight territories, the District of Columbia, and multiple health organizations and major hospital networks.</td>
<td>CDC</td>
<td>OEM</td>
</tr>
<tr>
<td><strong>Homeland Security Information Network (HSIN)</strong></td>
<td>HSIN is a comprehensive, nationally secure, and trusted web-based platform able to facilitate Sensitive But Unclassified information sharing and collaboration between Federal, State, local, tribal, private sector, and international partners. It interfaces with existing information sharing networks to support the diverse Communities of Interest engaged in preventing, protecting from, responding to, and recovering from all threats, hazards, and incidents. HSIN provides a collaborative environment that interoperates with mission area systems developed and managed by Federal, State, and local partners.</td>
<td>DHS</td>
<td>Centers, ORA, OEM</td>
</tr>
<tr>
<td><strong>Integrated Consortium of Laboratory Networks (ICLN)</strong></td>
<td>The ICLN was established to coordinate the Nation’s laboratory networks to improve the response to acts of terrorism and other events requiring an integrated laboratory response. One outcome of the ICLN is the creation of an Integrated Response Architecture that provides, among other things, event notifications and updates, preparedness alerts, and situational reports through a secure web portal.</td>
<td>DHS</td>
<td>ORA</td>
</tr>
<tr>
<td><strong>Laboratory Response Network (LRN)</strong></td>
<td>The LRN is a network of over 150 Federal, State, and local; military; food testing; environmental; veterinary; and international laboratories that are fully equipped to respond quickly to acts of chemical or biological terrorism, emerging infectious diseases, and other public health threats and emergencies.</td>
<td>CDC</td>
<td>ORA and CFSAN</td>
</tr>
<tr>
<td><strong>Manufacturer and User Facility Device Experience System (MAUDE)</strong></td>
<td>MAUDE data represents reports of adverse events involving medical devices. The data consists of voluntary reports, user facility reports, distributor reports, and manufacturer reports. The online search allows you to search CDRH database information on medical devices that may have malfunctioned or caused a death or serious injury.</td>
<td>FDA</td>
<td>CDRH</td>
</tr>
<tr>
<td><strong>MedWatch</strong></td>
<td>MedWatch, the FDA Safety Information and Adverse Event Reporting Program, provides important and timely clinical information about safety issues involving medical products, including prescription and over-the-counter drugs, biologics, medical and radiation-emitting devices, and special nutritional products. It allows healthcare professionals and consumers to report serious problems that they suspect are associated with the drugs and medical devices that they prescribe, dispense, or use.</td>
<td>FDA</td>
<td>CDER</td>
</tr>
<tr>
<td><strong>National Biosurveillance Integration Center (NBIC)</strong></td>
<td>The NBIC integrates, analyzes, and distributes key information about health and disease events to help ensure the Nation's responses are well-informed, save lives, and minimize economic impact.</td>
<td>DHS</td>
<td>OEM/OEO</td>
</tr>
<tr>
<td>System</td>
<td>Description</td>
<td>Sponsoring Agency</td>
<td>FDA POC</td>
</tr>
<tr>
<td>--------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>-------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>National Consumer Complaint System</td>
<td>FDA uses the National Consumer Complaint System to obtain complaint information from consumers about FDA-regulated products. Consumer Complaint Coordinators at FDA headquarters and District offices throughout the United States review complaints to identify product/problem trends and potential emerging public health issues.</td>
<td>FDA</td>
<td>OEM/OEO</td>
</tr>
<tr>
<td>New Drug Application (NDA) Field Alert Program</td>
<td>The NDA Field Alert Program is used to quickly identify drug products that pose potential safety threats. All drug manufacturers with approved NDAs and Abbreviated New Drug Applications (ANDAs) are required to submit Field Alert Reports to FDA if they find any significant problems with an approved drug.</td>
<td>FDA</td>
<td>CDER, ORA</td>
</tr>
<tr>
<td>The Pet Event Tracking Network (PETNet)</td>
<td>PetNet is a collaborative project between FDA and Federal and State government partners to provide a secure reporting/notification system, accessible by State and Federal government officials, which will allow for the exchange of information early in a disease outbreak that is associated with the consumption of adulterated pet food.</td>
<td>FDA</td>
<td>CVM</td>
</tr>
<tr>
<td>Potential Tobacco Product Violations Reporting (PTVR)</td>
<td>CTP uses PTVR to obtain complaint information from consumers and industry regarding potential tobacco-related violations. Submissions are reviewed by FDA’s CTP, Office of Compliance and Enforcement.</td>
<td>FDA</td>
<td>CTP</td>
</tr>
<tr>
<td>PulseNet</td>
<td>PulseNet is a national network of laboratories that performs DNA “fingerprinting” on foodborne bacteria. The network permits rapid comparison of patterns through an electronic database, helping to better detect the source of bacteria in foods. PulseNet helps public health authorities recognize when cases of foodborne illness are occurring at the same time in geographically separate locales.</td>
<td>CDC</td>
<td>CFSAN</td>
</tr>
<tr>
<td>Recall Enterprise System (RES)</td>
<td>RES is used by Center and District Office recall personnel to submit, update, classify, publicize, and terminate recalls. This online, searchable database allows users to enter real-time data on a recall event as it becomes available and track information and generate and disseminate reports of recall activities.</td>
<td>FDA</td>
<td>ORA</td>
</tr>
<tr>
<td>Safety Reporting Portal (alsoSRP or Reportable Food Registry [RFR])</td>
<td>RFR is an electronic portal for an owner, operator, or agent in charge of a domestic or foreign facility engaged in manufacturing, processing, packing, or holding food for consumption in the United States to file a report about a product that is suspected will cause serious adverse health consequences or death to humans or animals. Initial RFR reports are sent to EON-IMS.</td>
<td>FDA</td>
<td>OEM/OEO, CFSAN, CTP</td>
</tr>
<tr>
<td>Vaccine Adverse Event Reporting System (VAERS)</td>
<td>The VAERS is a national vaccine safety surveillance program to detect possible signals of adverse events associated with vaccines. VAERS collects and analyzes information from reports of adverse events (possible side effects) that occur after the administration of U.S.-licensed vaccines.</td>
<td>FDA, CDC</td>
<td>CBER</td>
</tr>
<tr>
<td>Veterinary Adverse Drug Experience (ADE) Reporting System</td>
<td>The ADE Reporting System provides early warning to FDA for adverse effects not detected during pre-market testing of FDA-approved animal drugs. Reports of adverse clinical events from veterinarians and animal owners are entered into a computerized database for use by scientists to make decisions about product safety, which may include changes to the label or other regulatory action.</td>
<td>FDA</td>
<td>CVM</td>
</tr>
<tr>
<td>Veterinary Laboratory Response Network (Vet-LRN)</td>
<td>The Vet-LRN enhances collaboration between Federal and State agencies and veterinary diagnostic laboratories and provides additional laboratory capacity and expertise. The network contributes to the protection of the human food supply as laboratory findings of contamination of animals feed or drugs could signal potential contamination to human food.</td>
<td>FDA</td>
<td>CVM</td>
</tr>
</tbody>
</table>
B.2.1.2 Consumer Product Protection

Prior to confirmation of an incident, FDA may employ a number of activities designed to deter intentional and unintentional acts against FDA-regulated products and protect consumers from potential public health hazards. Through heightened and targeted preventive measures at various points in the processing and distribution chains, and by exercising the systems and networks to be used during emergency response operations, FDA can protect the safety and security of regulated products and protect consumers from harm. Activities in support of this effort, conducted in cooperation and collaboration with applicable FDA organizational components, Federal and SLTT partners, industry, academia, foreign governments, and international organizations include, but are not limited to, the following:

- Implementing risk communications with government and industry partners.
- Prioritizing examination of food commodities based on potential for contamination (as determined by vulnerability assessments previously undertaken by FDA).
- Identifying entities handling specific FDA-regulated consumer products.
- Conducting targeted inspections and investigations and collecting samples.
- Readyling laboratory response and other scientific capabilities to analyze/test for chemical and microbiological agents.
- Tracing and intercepting specific imported and domestic products.
- Requesting the recall of drugs, biologics, medical devices, radiation-emitting devices, foods, or cosmetics that predictably could cause serious health problems or death (“Class I”) or that might cause a temporary health problem or pose only a slight threat of a serious nature (“Class II”).
- Requesting the recall of tobacco products containing a manufacturing or other defect not ordinarily contained in tobacco products on the market that would cause serious adverse health consequences or death.
- Providing additional guidance to food producers, processors, transporters, importers, retailers, food service establishments, and cosmetic processors on security measures to take to minimize the risk of tampering or other intentional contamination/adulteration and protect against the damaging effects of diseases and natural disasters.
- Working, as appropriate, with Federal and SLTT government agencies and international partners.
- Issuing warnings, alerts, advisories, and other advice to consumers, industry, foreign counterparts, and other international partners regarding food, drug, medical device, biological product, and tobacco product safety and animal health and soliciting feedback.

The FDA system for responding to the increased risk of an emergency or disaster impacting FDA-regulated products provides a significant increase in coverage, awareness, and preparedness through targeted preventive measures implemented by both headquarters and field offices and in coordination with other intergovernmental public health and medical partners. Collectively, these additional consumer

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7 Incident confirmation includes conclusive laboratory analysis, field investigations, epidemiological data, or reliable information from other government agencies that triggers emergency response operations.

8 Detailed FDA investigation/inspection procedures can be found at www.fda.gov/ora/inspect_ref. Compliance references are located at www.fda.gov/ora/compliance_ref.

9 Recalls of products that are unlikely to cause any adverse health reaction, but that violate FDA labeling or manufacturing laws (“Class III”), will not by themselves activate this EOP.
protection activities provide for more protected supply and distribution chains and a better prepared national network capable of responding to an identified threat or hazard.\textsuperscript{10}

\textbf{B.2.1.3 Medical Countermeasures}

FDA plays a vital and multifaceted role in securing the homeland through providing broad regulatory oversight, monitoring infrastructure, and responding to CBRN threats, whether intentional or naturally occurring with timely and appropriate medical countermeasures. FDA fosters the development of safe and effective medical countermeasures to mitigate the effects of such threats by actively engaging industry, academia, and working with our government partners. While all the FDA Centers and Offices have a role in the agency’s counterterrorism mission, the three product Centers (i.e., Center for Biologics Evaluation and Research [CBER], Center for Drug Evaluation and Research [CDER], and Center for Devices and Radiological Health [CDRH]), as well as ORA and various organizations within the Office of the Commissioner (OC), play critical roles in ensuring the safety, effectiveness, quality, and availability of medical countermeasures. The Office of Counterterrorism and Emerging Threats (OCET) provides strategic leadership and coordination for agency-wide counterterrorism and emerging threats activities.

In situations in which potentially useful medical countermeasures are available but not yet FDA-approved, licensed, or cleared for the particular use contemplated, OCET, in coordination with medical product Centers, can employ certain regulatory mechanisms to allow the use of these products during an emergency, as part of an expanded access Investigational New Drug (IND) application or Investigational Device Exemption (IDE), or under an EUA. In addition, there are other authorities to facilitate use of FDA-approved, licensed, and cleared medical countermeasures that may need to be used in ways that may render them in violation of the FD&C Act.

\textbf{B.2.1.4 Increased Surveillance of FDA-Regulated Medical Products}

Depending on the nature of an incident, FDA may increase its surveillance of FDA-regulated medical products used to prevent or mitigate adverse health effects related to the actual or potential incident. Such increased surveillance could be a result of conditions such as the use of a product by a larger or different population, the use of products for other than their labeled indication, or concerns about the potential development of resistance to a product such as antivirals or antibiotics.

\textbf{B.2.2 Response}

Under the NPG, the response mission area includes those capabilities necessary to save lives, protect property and the environment, and meet basic human needs after an incident has occurred.

The “response” phase of FDA emergency operations includes those immediate and sustained actions to ensure the safety, efficacy, and availability of FDA-regulated products and to protect the public’s health throughout the duration of an incident. This phase generally involves the following four elements

1. \textbf{Gain and Maintain Situational Awareness}
   - \textit{Alert and Notification}. Receiving information or intelligence confirming the development or occurrence of an incident and issuing notifications to agency stakeholders and partners.
   - \textit{Assessment and Monitoring}. Performing initial and ongoing situation analysis, monitoring, and reporting.

\textsuperscript{10} For more information on FDA protection measures as part of the Nation’s larger critical infrastructure protection efforts, refer to the National Infrastructure Protection Plan’s sector-specific plan for food and agriculture located at https://www.dhs.gov/sites/default/files/publications/nipp-ssp-food-ag-2015-508.pdf.
2. **Activation and Deployment of Resources and Capabilities.** Directing and mobilizing FDA headquarters and field resources and capabilities, including Emergency Operation Center (EOC) operational-level changes and implementation of the ICS in the field and/or headquarters.

3. **Coordination of Response Actions.** Conducting rapid and synchronized emergency response activities at headquarters and field locations.

4. **Demobilization.** Standing down FDA emergency response resources (i.e., personnel, facilities, equipment, and other materials support) and returning them to their original location and status.

![Figure B-1. FDA Emergency Response Elements](image)

**B.2.2.1 Gain and Maintain Situational Awareness**

**B.2.2.1.1 Alert and Notification**

FDA may be alerted to a threat, hazard, or other significant event through a variety of means, including the surveillance systems discussed in Table B-1 and/or directly from external Federal agencies (e.g., the HHS Secretary’s Operations Center [SOC], Centers for Disease Control and Prevention [CDC], U.S. Department of Agriculture [USDA], DHS, and/or the Federal Bureau of Investigation [FBI]); international and SLTT public health agencies; industry; consumers; news media; and internal FDA organizational components. Any or all agency organizational components may receive word of a problem depending on the event’s scope and magnitude. Recipients of such information must handle it properly to ensure FDA is able to track and evaluate the situation. Failure to address a report in a timely fashion may delay emergency response and potentially exacerbate the incident.

Depending on the scale of the incident, headquarters coordination of an incident will be managed by OEO Emergency Coordinators or through activation of an IMG (in either case, coordination may be conducted from FDA’s Emergency Operations Center (EOC) or other locations as circumstances may require.

The EOC is staffed by OEM/OEO staff members who work with the Centers and Offices to review information that may indicate potential or actual incidents, as well as quickly detect similarities in seemingly unrelated reports, and ensure that incident reports are treated in a consistent manner across FDA. The EOC also provides or acquires from a variety of sources the resources and capabilities necessary for FDA headquarters-level and interagency coordination that may be necessary when emergencies or disasters cross geographic and/or jurisdictional boundaries.

All reports of large-scale natural and manmade emergencies and significant alleged or actual adverse effects associated with FDA-regulated products require prompt reporting to OEM/OEO. Upon receipt of an alert or warning, the OEM/OEO staff will notify internal and external partners via a predeveloped, prioritized notification list specific to each particular emergency.
Certain types of incidents typically, but not necessarily always, should be reported to OEM/OEO as potential emergencies. These include incidents that:

- Involve life-threatening adverse events or deaths possibly related to use of drugs, biologics, medical devices, radiation-emitting devices, the food supply, or cosmetics.
- Involve life-threatening adverse events or deaths possibly related to use of an adulterated tobacco product containing manufacturing or other defects not ordinarily contained in tobacco products on the market.
- Involve a contaminant or pathogen associated with FDA-regulated products.
- Involve product distribution and/or injury, illness, or death across multiple States or geographic regions.
- Involve a considerable number of producers or manufacturers or a firm of considerable size and extensive product distribution.
- Arise in circumstances in which it is difficult to quickly identify the source of the problem.
- Involve multiple FDA organizational components.
- Involve natural and manmade disasters.
- Involve terrorist and/or criminal activities with potential to impact FDA staff, facilities, or regulated products.

**B.2.2.1.2 Assessment and Monitoring**

Once FDA officials receive word of an impending, ongoing, or resurgent emergency situation, they must gather enough information to determine the validity and extent of the threat or hazard and the agency’s roles and responsibilities within the situation. FDA headquarters and field organizational components work collaboratively with one another and with other government agencies and industry to rapidly evaluate the situation and make these determinations.

OEM/OEO is responsible for coordinating with applicable FDA headquarters, field units and external partners to formulate an initial incident assessment. Results of this assessment may include a variety of situation reports (SitReps) and status reports, healthcare records, research and analyses, maps, and surveillance information. This critical information is passed through established reporting channels to allow agency decision-makers to develop situational awareness and establish a “common operating picture.” Using OEM’s GIS capabilities, an online or paper map-based common operating picture may be used to build and maintain situational awareness. The map-based common operating picture can visually depict response features with geographical reference in addition to the text information. Based on initial analysis of the threat or hazard, FDA will take steps to monitor the situation, identify and prioritize requirements, and/or activate available resources and capabilities. Figure B-2 depicts the methodology agency officials use to assess an incident and determine whether additional response activities are warranted.

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11 At the time of the initial notification, an FDA organizational component or another government agency may already have completed a preliminary assessment of the emergency.
Figure B-2. FDA Emergency Response Decision Tree

Determining Regulatory Authority. It is important to note that a reported incident may or may not be subject to FDA authority. Therefore, one of the first discussions to take place will include a determination of whether FDA has regulatory authority. Some of the criteria potentially used in the assessment include:

- Whether an FDA-regulated product is implicated as the cause of the incident.
- Whether it is confirmed that FDA-regulated products have been or may be affected.
- Whether medical countermeasures are expected to be used in response.
- Whether interstate commerce of FDA-regulated products, as defined under the various statutes FDA enforces, is involved.
- Whether Federal or SLTT agencies and/or foreign governments have formally requested FDA assistance.

Once FDA determines that it has regulatory authority, it may manage the incident internally or function in a supporting role. Although the agency may make an initial determination regarding authority and jurisdiction, it may continue to review and reconsider regulatory authority throughout an incident as it evolves, expands, or contracts.

B.2.2.2 Activation and Deployment of Resources and Capabilities

As the agency’s central coordination point for emergencies, the FDA EOC operating status generally falls into one of three levels, depending on the severity or potential consequences of an incident or perceived threat (Table B-2). Investigation or confirmation of an incident with potential or confirmed public health impact may require activating a change in the EOC operational level from a readiness state of Operational Level 3 to Operational Level 2 or 1, depending on the scope of the incident. Center and District Office Situation Rooms, Command Centers, and Emergency Response Coordination Centers may follow the EOC operating levels or activate according to their individual emergency plans and procedures as appropriate.

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12 Refer to Appendix A and Appendix B of this EOP for detailed information on how FDA supports other Federal agencies under the NRF.
Table B-2. FDA Incident Levels and EOC Operation Levels

<table>
<thead>
<tr>
<th>Operational Level</th>
<th>Operating Status</th>
<th>Criteria</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>Routine awareness and response operations</td>
<td>Incident(s) involving FDA-regulated product(s) that, from a response and coordination viewpoint, are limited in complexity and scope such that the resources needed for response and/or coordination do not significantly impact normal operations for response/coordination. OEM and OEM/OEO coordinate with ORA and Center emergency coordinators. Generally, FDA’s response and coordination actions are sufficient to respond effectively to the incident with no or limited internal and/or external coordination necessary.</td>
<td>Single-person illness, injury, or consumer complaint with no or very few similar complaints in FDA systems</td>
</tr>
<tr>
<td></td>
<td>Normal staffing</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>Regular work hours</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>Readiness state</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Increased response operations</td>
<td>Incident(s) involving FDA-regulated products(s) that, from a response and coordination viewpoint, are of moderate complexity and scope such that resources needed for an effective response and/or coordination are beyond those ordinarily provided during the routine workday and/or beyond average staffing levels. The incident may require periodic or ongoing coordination with other external entities, in which case FDA may or may not be the “lead” responding agency for the incident. If additional staff and resources are not provided, response and coordination efforts would be somewhat adversely impacted, most likely in terms of response timeliness.</td>
<td>CDC report of microbial contamination in pharmacy-compounded drugs Ongoing foodborne illness outbreak involving one or more States (e.g., Salmonella Montevideo outbreak in 2009-2010 and Salmonella Enteriditis outbreak in 2010) H1N1 response in 2009 NSSEs National-level or full-scale agency exercises</td>
</tr>
<tr>
<td></td>
<td>Augmentation of EOC staffing may be necessary</td>
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<tr>
<td></td>
<td>Extended work hours as needed</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Center Situation Room activation may be necessary</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Establishment of an Incident Management Group (IMG) may be necessary (EOC serves as resource to IMG)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Escalated response operations</td>
<td>Incident(s) involving FDA-regulated product(s) that, from a response and coordination viewpoint, are of substantial complexity and scope such that the resources needed for an effective response and/or coordination exceed the Level 2 capability. Staffing levels, hours, and operations will likely be substantial to mount an effective response (to prevent or mitigate public health impact). Incidents may require significant coordination with external entities to optimize response. FDA may be the “lead” responding agency and/or may have a substantial support role to provide to another “lead” agency. Both field and headquarters response operations and/or coordination activities may require the establishment of formal organizational restructuring (e.g., ICS) to respond optimally.</td>
<td>Major natural disaster involving multiple States Ongoing foodborne illness outbreak or food contamination involving multiple States and increased complexity in product tracing involving multiple agencies Unknown contaminated drug product with serious adverse event Contaminated human food, pet food, or animal feed Terrorist event</td>
</tr>
<tr>
<td></td>
<td>Additional personnel (support and subject matter experts [SMEs])</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Extended work hours likely, 24-hour operations possible (8- to 12-hour shifts)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Center Situation Room provides EOC daily SitReps</td>
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</tbody>
</table>

Depending on the anticipated or actual size and complexity of an incident, activation of FDA emergency management resources may be required. The EOC, staffed by OEM and OEM/OEO (typically Operational Level 3), coordinates with ORA’s headquarters and field components and the appropriate Center(s) in the review and analysis of information about threats and hazards and assists in the early recognition of emergencies, outbreaks, natural disasters, and terrorism or other criminal acts that may affect an FDA-regulated industry or product. This agency-wide coordination involves information collection and distribution, triaging complaints and alerts, issuing mission assignments (MAs) to organizational components, identifying and tracking needed resources, and communicating with external partners as they
request technical and material support. Additionally, the FDA EOC serves as the central link for coordination between FDA components and the HHS SOC, the CDC Director’s EOC, USDA’s EOC and/or Food Safety and Inspection Service (FSIS) Situation Room, and other Federal departments/agencies as appropriate.

**B.2.2.2.1 Authority to Change Operational Levels**

The authority to change the operational level of FDA’s EOC resides with the OEM Director, OEM/OEO Director, or his/her designee. The OEM or OEM/OEO Director will coordinate changes in operational levels with Office of Operations (OO)/Office of Security and Emergency Management (OSEM) leadership as needed. The Commissioner or the Secretary or designee also may request a change in the operational level of the FDA EOC. Other agency emergency coordination units, such as Centers, shall activate their Situation Rooms at the discretion of senior management, such as the Center Director. District offices will establish Incident Management Teams (IMTs) at the discretion of the District Director (DD) or Program Director (PD). FDA’s EOC (OEM/OEO) shall be made aware when a Situation Room or IMT is activated in response to an incident. An increase in the EOC operational level may lead to the establishment of an Incident Management Group (IMG) to bring together additional resources and expertise to assume overall coordination for the response. For FDA field response, see Section B.2.2.3.2.

**B.2.2.2.2 Conditions for Operational Level Change**

Any of the following conditions could affect the operational levels of the EOC or activation of the FDA IMG:

- FDA has received credible intelligence or other confirmed information that an FDA-regulated product is the specific target of terrorism or other serious criminal activity.
- An incident involves a single or multiple FDA-regulated product(s) covering a large geographic or population area (more than one State) and requires the coordination of multiple agency organizational components.
- Two or more significant events have occurred at the same time.
- FDA-regulated products or facilities have been impacted by a natural or manmade disaster.
- A natural or manmade incident has triggered activation of the NRF, in whole or in part, and State or Federal agencies have formally requested assistance from FDA.
- Deliberate or accidental contamination of the human or animal food, medical products, or tobacco products supply has caused widespread illness, injury, or death to consumers.
- A widespread epidemic or pandemic disease outbreak or other public health emergency for which FDA has a designated support responsibility has occurred or is highly probable.
- DHS has raised the NTAS level to **Elevated** (“Warns of credible terrorist threat against the United States”) or to **Imminent** (“Warns of credible, specific, and impending terrorist threat against the United States”).

The FDA EOC will notify appropriate agency organizational components, the HHS SOC, and other key Federal partners when the EOC operational level has been changed or an IMG has been activated in response to an emergency. This activation announcement may include a status report explaining the rationale for the decision to change the EOC operational level or activate an IMG, a description of

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13 For detailed information on emergency alert and notification procedures, refer to Section E, “Communications and Information Management,” of this EOP.
operating hours and staffing levels, and situation reporting requirements. After activation procedures commence, FDA may request additional resources and capabilities to augment emergency operations. This includes prioritizing and clearly communicating specific incident requirements to mobilize and deploy emergency response staff to both internal and external locations. Requests for staffing will be made in writing by the OEM Director, OEM/OEO Director, or someone higher in the OC to Center and Office lead emergency coordinators or other senior officials within the organization. Representatives from specific headquarters Centers and Offices may be directed to relocate to the FDA EOC, other designated FDA locations, or other Federal departments or agencies. Interagency partners may also send representatives to the FDA EOC or other agency organizational components to perform liaison roles under existing memorandums of understanding (MOUs). In addition, FDA field organizations, such as, District offices, branches, and resident posts may send personnel and specialized equipment to the FDA EOC, incident site(s), laboratories, or other designated locations to provide subject matter expertise and advice, conduct investigations and inspections, collect and analyze samples, and/or detain or destroy contaminated product(s).

EOC Operations During Voluntary Isolation. The physical staffing of the IMG may take place within the location of the EOC or another designated location if needed. However, various conditions associated with a pandemic (i.e., quarantine, social distancing practices, and the need for staff to care for seriously ill family/friends), natural disaster, or other event impacting EOC members’ ability to physically staff the EOC may make it necessary at times for IMG or EOC assigned staff personnel to carry out their duties from a remote location, such as their residence. The authority to operate the FDA EOC under “virtual” conditions resides with the OEM Director, OEM/OEO Director, or his/her designee.

B.2.2.3 Coordination of Response Actions

FDA personnel must perform a number of tasks and functions when responding to an incident. Some of these basic tasks may include performing initial and ongoing planning, managing and performing MAs, coordinating team response operations and resources, performing research and decision-making, and documenting and reporting information. Although FDA personnel accomplish most of these basic tasks over the course of the incident, their order of execution, the exact activities performed, and the individual or organization responsible will vary depending on the type and severity of the threat or hazard. Each involved FDA organizational component will be responsible for performing certain emergency actions based on its assigned mission and functions.

FDA organizes emergency response operations in accordance with the concepts, principles, and terminology of the ICS, as defined within NIMS. Incident Command and subordinate Planning, Operations, Logistics, and Finance/Administration functions lay the foundation for the agency’s implementation of the ICS during all-hazards response, at headquarters and field levels, and across geographic divides. The inherent design of this system enables rapid, scalable, and flexible agency-wide emergency management activities.

This section describes each of the components of a coordinated FDA emergency response. The agency’s response strategy, while containing certain core activities, must be tailored to the specific requirements of each incident. (Refer to the Incident Annexes of this EOP for detailed information on FDA functions, activities, and organizational constructs for individual hazards.)

14 For more information on MOUs, refer to the FDA EOP Appendix I.
15 Refer to Section D, “Direction, Control, and Coordination,” of this EOP for more information on agency emergency position assignments.
16 Refer to Section C, “Organization and Assignment of Responsibilities,” of this EOP for more information on individual Center/Office roles and responsibilities.
17 For more information on the ICS, refer to www.training.fema.gov/EMIWeb/IS/ICSResource/index.htm.
B.2.2.3.1 Initial and Ongoing Response Planning

FDA conducts planning to organize its response structure, identify objectives and performance metrics, and coordinate the delivery of resource support. Planning is ongoing throughout the lifecycle of any incident and is adjusted as necessary to meet changing demands. It involves a blend of prescribed actions drawn from existing FDA emergency plans and procedural documents (e.g., hazard-specific incident annexes, SOPs, operations manuals, field guides) and real-time determination of the necessary course(s) of action. An agency Incident Action Plan (IAP) may be developed based on these actions to provide operational and tactical direction to FDA emergency personnel when responding.

The IAP is generally crafted by holding regular planning meetings throughout the incident, both in-person and via conference calls, with responsible headquarters and field units. Planning meetings seek to accomplish the following goals:

- Gathering, recording, analyzing, and distributing incident information in a manner that will facilitate: (1) increased situational awareness of the magnitude, complexity, and potential impact of the incident, emergency, or crisis; and (2) the ability to determine the resources required to resolve the situation.
- Formulating and prioritizing measurable incident objectives and identifying an appropriate response strategy that conforms to the legal obligations and management objectives of all FDA organizational components and/or external agencies involved.
- Determining the tactical direction, reporting mechanisms, timeframes, and specific resource requirements (i.e., personnel, specialized equipment, facilities, training and expert knowledge, and funding) for implementing selected strategies during the response period.

Through these planning meetings, FDA can coordinate the execution and evaluation of emergency activities, promote and maintain situational awareness (a “common operating picture”), track the progress of ongoing initiatives, and modify plans and procedures based on new and emerging information. For incidents involving both a headquarters (IMG) and field (IMT) response, IAPs will generally be developed by field units, while an Incident Coordination Plan (ICP) may be developed by the IMG that outlines incident objectives from a headquarters perspective (e.g., support of field operations).

OEM/OEO can establish an Intelligence Unit (within the IMG Planning Section) to provide staff support if necessary. This unit can provide: 1) coordination of information collection and initial response activities of Centers, Offices, and Districts; 2) collection and assessment of surveillance data; 3) initial development of forward planning; and 4) advice to the OEM Director (refer to Section D.2.2.2.2, “Intelligence Unit”).

B.2.2.3.2 FDA Field Response

For the majority of incidents involving FDA-regulated products, one or more FDA field offices may be involved in the response. The Districts in which the event is occurring (i.e., the physical location where people have been affected) will obtain necessary information for FDA to confirm the health hazard. In addition, the field offices will determine, plan, and conduct tasks and assignments over the course of the response.

Incident Management Team. A decision can be made to respond by implementing ICS at the field level, which involves the activation of one or more IMTs. An IMT utilizes an incident command structure made up of the Command, General Staff members, and appropriate functional units. Criteria for activating an IMT will be based on the conditions described in Section A.3, “Scope and Applicability.” The authority to mobilize an IMT resides with ORA senior field officials, in consultation with senior ORA HQ officials. Ideally, IMT activation should be coordinated with OEM or with the IMG if activated, but, at a minimum,
these organizations should be notified whenever an IMT is being activated. An IMT may also be mobilized at the request of OEM or an IMG.

The IMT will be responsible for tactical operations (i.e., perform investigations/inspections, collect samples, and/or detain or destroy contaminated product) in accordance with the IAP it develops. In addition, as the IMT conducts its follow-up efforts, field offices will communicate investigational/inspectional findings of other potentially affected establishments with the firms’ home Districts/Programs and FDA headquarters emergency response entities described in the following section. Every time an IMT is activated, a Program Executive Group (PEG) will be activated as well to provide program-specific overarching strategy and policy guidance at the district/geographic level (see Section D.1, “Field Incident Command”).

The IMT is led by a designated Incident Commander (IC) (see Section D.1, “Field Incident Command,” for organizational elements). The field IC reports directly to the PEG, or FDA Area Command, if activated, and provides situational awareness to FDA’s OEM/OEO or an IMG (see Section B.2.2.3.3, “FDA Headquarters Response,” for IMG description). The IC coordinates activities with the IMG (when activated). Communications are conducted through teleconference calls, emails, and/or SitReps. The IMT is responsible for managing the District’s/Program’s emergency response effort, coordinating actions and resources needed to follow up on the incident, and channeling all necessary communications to/from deployed field elements. Depending upon the nature and severity of the incident, additional field IMTs may be established to support geographically distant or functionally different emergency response operations.

Generally, the field IMT’s location could be set up at any of the FDA field offices near the site of investigational or inspection activities because of the accessibility to records and the availability of dedicated communications/information systems equipment. However, if the incident has destroyed or otherwise negatively impacted these offices, consideration may be given to relocating the Incident Command Post (ICP) to an alternate facility (such as a predesignated District Office continuity of operations (COOP) site or other FDA District Offices). A notification for reallocation of field staff between or among District Offices shall be directed to ORA Assistant Commissioners (ACOs) in coordination with the Associate Commissioner for Regulatory Affairs (ACRA). As circumstances warrant, an IMT could be located at a location other than an established FDA Office, such as collocation with an HHS Incident Response Coordination Team (IRCT), FEMA Joint Field Office (JFO), hotel, or a facility near an establishment being inspected, as appropriate.

### B.2.2.3.3 FDA Headquarters Response

Depending on the nature and complexity of FDA’s response to an incident, an intra-agency approach may be used to coordinate resources and incident-related information and to support incident management and agency policies. Depending on the scale of the incident, headquarters coordination will be managed by OEO Emergency Coordinators or through activation of an IMG; in either case, coordination may be conducted from FDA’s Emergency Operations Center (EOC) or other locations as circumstances may require.

- **IMG.** An IMG is typically established when an incident involves multiple FDA organizational components or involves (or is anticipated to involve) complex incident management/coordination. The authority to establish FDA’s incident management structure, IMG, resides with the OEM Director, OEM/OEO Director, or his/her designee, in consultation with senior Center/Office and ORA leadership, as appropriate. It is staffed by agency headquarters representatives (i.e., from the OC, OEM, and/or the FDA product Centers) and, as appropriate, FDA field staff and/or external agency liaisons. The IMG staffing is organized consistent with the principles of NIMS. An Agency Incident Coordinator (AIC) is designated to oversee the IMG, including development and
implementation of a headquarters ICP and the coordination of FDA headquarters resources and capabilities to respond to the event, as needed. Subordinate Command and General Staff positions, comprising representatives from headquarters Centers and Offices, are also established to assist the AIC in executing emergency functions under the specific conditions at hand.\textsuperscript{18} The AIC reports directly to the Commissioner or the Commissioner’s designee, an Agency Executive Group (AEG), or the FDA Executive Committee (provides oversight and direction on cross-cutting operational issues, strategic challenges, and scientific policy issues for FDA). An AEG or Executive Committee may provide guidance on policy issues and resolve issues involving competing resources. For a further description of the AEG and/or the Executive Committee, see Section D, “Direction, Control, and Coordination.”\textsuperscript{19}

The scope of the incident may call for activation of an internal FDA Joint Information Center (JIC). This is a collocated group of communication specialists from Centers and Offices involved in the event that are designated to handle public information needs. For a further description of the FDA JIC, see Section D.2.2.4.1, “Units.”

- **EOC.** The FDA EOC is the physical location at which the coordination of information and resources to support incident management activities normally takes place.

**FDA EOC.** The FDA EOC serves as the agency-wide focal point for emergency operations coordination and dissemination of information, whether staffed by OEM/OEO personnel or an IMG. The FDA EOC staff monitors ongoing events, processes complaints and alerts, issues MAs to FDA organizational components, coordinates overall agency emergency management operations, and communicates with interagency partners to provide technical and material support.\textsuperscript{20} The FDA EOC staff facilitates contact between applicable headquarters and field emergency personnel and provides frequent and formalized communications/reporting to senior agency officials and organizational components, as well as ASPR and the HHS SOC and other external partners, regarding the status of FDA emergency response activities.

The FDA EOC provides a physical central location from which headquarters personnel can provide agency-wide coordination and executive decision-making in support of the incident response. The FDA EOC staff does not command or control the agency’s response, but carries out the coordination function for complex incidents or multiple incidents occurring simultaneously through:

- **Information Collection, Evaluation, and Dissemination.** Collecting, analyzing, interpreting, and distributing information from/to various internal and external sources.

- **Priority Setting.** Ensuring that agency response systems are interconnected and complementary, reinforcing interoperability among the various organizational components, making the response more efficient and effective by coordinating available resources, and making decisions based on agreed-upon policies and procedures.

- **Resource Coordination.** Identifying and acquiring needed resources and allocating existing or known resources.

- **Communications Facilitation.** Establishing and maintaining intra- and interagency interoperable communications.

\textsuperscript{18} For detailed information on how FDA organizes and staffs emergency response operations, refer to Section D, “Direction, Control, and Coordination,” of this EOP.

\textsuperscript{19} The use of the terms “Incident Management Group” and “Agency Executive Group” should be reserved for use only as defined in the FDA EOP. Use of ICS principles across FDA is encouraged; however, the terms IMG and AEG only apply to the incident structure activated by OEM/OEO and are used as described above.

\textsuperscript{20} The composition of FDA EOC staff will change depending on specific incident requirements.
**Centers.** The FDA product Centers are responsible for scientific evaluations and for programmatic decisions and policy development within their respective program areas. Their professional staff includes clinical, scientific, and manufacturing process, policy, as well as medical countermeasure and emergency coordination experts. This expertise (analytical, laboratory, sampling procedures, subject matter expertise, and industry knowledge) is routinely utilized for critical consultation and decision-making when incidents occur.

Centers may maintain a Situation Room or Coordination Center consisting of Center-selected subject matter experts (SMEs) to be led by a Center Emergency Coordinator (CEC). The CEC reports to the Center Director (or other senior leader) and coordinates directly with OEM/OEO Emergency Coordinators or an IMG. The CEC serves as the focal point for internal and external Center communications during emergencies and disasters and is responsible for providing regular status and SitReps to the FDA EOC as requested. FDA Centers participate in agency emergency operations when the response includes, or may include, regulatory activities or products under their jurisdiction. During some large-scale or catastrophic incidents, CECs or other Center personnel may be directed to report, or connect virtually, to the FDA EOC and an OEM/OEO Emergency Coordinator to support initial or prolonged emergency operations as part of an IMG. For further information about the Center’s responsibilities during an emergency, please see Section C.3, “Office of Foods and Veterinary Medicine,” and Section C.4, “Office of Medical Products and Tobacco” of this EOP.

**ORA.** Any information received by ORA regarding the incident or ongoing field activities will be shared and/or discussed, as appropriate, with staff operating from the FDA EOC and the Commissioner or the Commissioner’s designee as appropriate. Additionally, ORA, working with the responsible Centers, will develop, issue, and approve any new or revised regulatory policy required during the event.

For further information about ORA’s responsibilities during an emergency, please see Section C.5, “Office of Global Regulatory and Policy,” of this EOP.

**B.2.2.3.4 Mission Assignments in Support of State and Local Response**

Requests for support from State and local agencies can be made directly to the agency or, in the case of a declaration of a major disaster or emergency under the Stafford Act, through FEMA. In order to assist SLTT agencies with such requests, FDA has developed Pre-Scripted Mission Assignments (PSMAs), which can be used to expedite the submission of requests when a major disaster or emergency has been declared. These requests include laboratory analysis, investigational, sampling collection, subject matter expertise, and training related to drugs, biologics, medical devices, radiation-emitting devices, tobacco, the food supply, or cosmetics. See Appendix C, “Processing Mission Assignment Sub-Taskings,” for the list of the PSMAs and the procedures on executing MA sub-taskings.

**B.2.2.3.5 Use of Investigational Medical Products, including Medical Countermeasures**

When a countermeasure has not yet been approved, licensed, or cleared because the safety, effectiveness, and/or quality have not been fully established, FDA may permit its use under an IND application or IDE through expanded access mechanisms or under an Emergency Use Authorization (EUA).

Expanded access (sometimes called “compassionate use”) is the use of investigational drugs, biologics or medical devices outside the clinical trial setting to diagnose, monitor, or treat patients with serious diseases or conditions for which there are no comparable or satisfactory approved therapy options available. Whenever possible, accessing such an investigational product through a clinical trial is preferable because clinical trials can generate data that may lead to the approval of products and, consequently, to wider availability. However, when patient enrollment in a clinical trial is not possible

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21 FD&C Act, Section 561, (b) and (c): Expanded Access to Unapproved Therapies and Diagnostics.
(e.g., a patient is not eligible for any ongoing clinical trials, or there are no ongoing clinical trials), patients may be able to receive the product, when appropriate, through expanded access.

For certain events, the Commissioner may issue an EUA to allow the use of an unapproved medical product or the unapproved use of an approved medical product. Before FDA may issue an EUA, the HHS Secretary must declare that circumstances exist justifying the authorization, based on one of the following actions:

- A determination by the Secretary of Homeland Security that there is a domestic emergency, or a significant potential for a domestic emergency, involving a heightened risk of attack with a CBRN agent(s);
- A determination by the Secretary of Defense that there is a military emergency, or a significant potential for a military emergency, involving a heightened risk to United States military forces of attack with a CBRN agent(s);
- A determination by the Secretary of HHS that there is a public health emergency, or a significant potential for a public health emergency, that affects, or has a significant potential to affect, national security or the health and security of United States citizens living abroad, and that involves a CBRN agent or agents, or a disease or condition that may be attributable to such agent(s);
- The identification of a material threat, by the Secretary of Homeland Security pursuant to section 319F-2 of the Public Health Service (PHS) Act, that is sufficient to affect national security or the health and security of United States citizens living abroad.22,23

Although FDA clearly defines most medical countermeasure regulatory jurisdictional responsibilities, several jurisdictional situations require explanation. CBER regulates all vaccines and blood derivatives for human use, and the USDA regulates those for veterinary use. CDER regulates all drugs for human use, and the Center for Veterinary Medicine (CVM) regulates all drugs for veterinary use. CDRH regulates the majority of clinical diagnostics; however, CBER regulates a specialized subset of in-vitro diagnostics (IVD) involved in the safety of the blood/tissue supply. Therapeutic biologics are regulated by multiple Centers, and questions should be addressed to the appropriate Center. In instances where Center-specific jurisdiction within FDA is unclear, the FDA Office of Combination Products (OCP) will decide which FDA Center possesses regulatory jurisdiction.

Responsible OC Offices (including OCET) and FDA Centers and Offices shall perform the following general functions related to medical countermeasures:

- Monitoring and updating the status of the product regulatory category, availability, and relevant manufacturing information.
- Facilitating the availability of safe and effective drugs, therapeutic biologics, vaccines, cellular- and tissue-based therapies, diagnostic devices, and other biological products to be used as countermeasures.
- Collaborating with Federal and SLTT public health agencies and regulated industry regarding product stockpile issues, including labeling, appropriate usage, product performance, monitoring, use in special populations, and other evolving issues.

22 The most current FDA guidance on EUAs can be found at www.fda.gov/RegulatoryInformation/Guidances/ucm125127.htm.
23 FD&C Act, Section 564, Authorization for Medical Products for Use in Emergencies.
• Conducting prospective discussions about emergency use of products and the appropriate regulatory mechanism for their use and the submission of data (i.e., EUA, IND, IDE, or Master File) on the product.

• Ensuring product recipients will be provided information (1) adequate to obtain informed consent (for IND and IDE products), or (2) as required by the EUA (e.g., that the product is under EUA and that the known and potential risks and benefits of taking or refusing the product are adequately communicated).

• Providing advice on identifying potential contaminants of drugs and therapeutic biological products and advising how to test for quality and purity.

B.2.2.4 Demobilization

As the need for full-time incident response coordination wanes, FDA will assess the situation to determine whether to continue or terminate emergency operations.24 This decision includes identifying whether consumer product safety and security and public health protection objectives were achieved and an orderly, safe, and efficient return of FDA resources (i.e., activated or deployed personnel, facilities, and equipment) to their original locations and/or operating status is warranted. In making this decision, FDA coordinates with and seeks input from appropriate experts and stakeholders. As a result, FDA organizational components may resume normal business operations and transition to short- and long-term recovery. Demobilization of an IMG will be announced through written correspondence from the AIC to all IMG members, generally as part of a written demobilization plan developed by the IMG Planning Section. The FDA EOC, or IMG, if activated, should be notified whenever an IMT has been demobilized.

Depending on the incident, resurgence may occur, and any period of transition back to normal operations could be disrupted. When this occurs, the agency will begin a new “notification and alert” period for emergency response and a change in EOC operational levels may occur.

Transition back to normal operations, after initiating and conducting an emergency response, will occur in stages and will correspond to the recovery of affected communities, FDA-regulated firms, and the Nation as a whole.

B.2.3 Recovery

Under the NPG, the recovery mission area includes those capabilities necessary to assist communities affected by an incident in recovering effectively.

The “recovery” phase of FDA emergency operations includes short- and long-term actions to ensure and restore the safety, efficacy, and availability of FDA-regulated products and to protect the public’s health. To accomplish this, FDA supports HHS in the execution of the NDRF and the two Recovery Support Functions (RSFs)—Health and Social Services and Infrastructure Systems—that require HHS coordination or support.

As Federal ESF operations conclude, requests for more detailed assessments, technical assistance, and longer term coordination fall under the recovery mission. Subject matter expertise and technical assistance can be provided to address issues that impact FDA-regulated industries. FDA and Federal supporting agencies will work with SLTT entities to provide a coordinated and consistent approach to recover from an incident. FDA will collaborate and maintain communication with other government agencies, industry, the public, and the media particularly concerning any long-term affects associated with the event.

24 Only the Commissioner, his/her authorized designee, or the AIC may formally terminate agency-wide emergency response operations. However, Center Directors, DDs, and PDs, through their respective emergency coordinators, are authorized to demobilize emergency personnel within their organizational components (e.g., IMTs).
Recovery objectives that support Federal and SLTT agencies for each RSF are outlined below:

- **Health and Social Services**
  - Inspect or investigate FDA-regulated facilities and collect and analyze samples of FDA-regulated products for the overall recovery operation as needed.
  - Conduct prolonged inspection, sample collection, and laboratory analysis activities in support of SLTT agencies that are specifically related to a disaster but deemed ineligible under the Stafford Act (e.g., inspections of retail food establishments and pharmacies).
  - Manage a potentially large number of samples of FDA-regulated products, including long-term sampling associated with the health effects of natural and manmade disasters.
  - Oversee product destruction and product reconditioning or rendering as appropriate.

- **Infrastructure Systems**
  - Identify key pharmaceutical and medical device facilities and determine if any supplies of drugs, biologics, or medical equipment have been or will be critically impacted.
  - Facilitate the availability of critically needed medical products by identifying alternate products or sources of products.
  - Identify key facilities that manufacture/process, pack, or hold food for human or animal consumption and determine if any are critically impacted.
  - Provide clear and consistent information to assist industry in understanding and complying with regulations in a post-disaster environment.

Dependent on the recovery objectives and the level of coordination needed, the FDA IMG will determine if FDA EOC or FDA JIC resources are needed to support the functioning of the IMG and recovery efforts.

As the need for recovery coordination wanes, FDA will assess whether consumer product safety and security and public health protection objectives were achieved. The transition back to a steady state operation will occur in stages and will correspond to the reestablishment of public confidence in FDA-regulated products. In regard to the recovery of FDA-regulated industry, it may be more appropriate that long-term recovery activities requiring FDA involvement be coordinated by agency components other than OEM/OEO or an IMG.


**B.2.3.1 Evaluation of Response**

Following any emergency or disaster, FDA organizational components will discuss how the agency handled the incident and seek input from appropriate experts and stakeholders. Participants will analyze emergency actions and responsibilities, resolve any deficiencies, mitigate consequences, and anticipate and address any long-term affects of the incident. In addition, organizational policies, plans, and procedures will be updated as needed, incorporating lessons learned and best practices captured during the event.

Following the termination of response activities that involved an IMG, OEM works with stakeholders to conduct a lessons learned analysis to identify the following:

- Strengths and weaknesses of key response activities performed during the incident, including agency measures to protect public health.
- Resource needs (e.g., personnel, equipment, training).
- Improvements to emergency response plans and procedures.
- Strengths and weaknesses of agency communications, which could include intra-agency and interagency communications as well as communications with specific stakeholder groups, such as industry, consumers, and the news media.
- Needed modifications toward regulatory policy, laboratory and field operations, and research activities.
- Any other needed improvements to overall preparedness.

As an important part of this process, it is generally advisable that when an IMG has been activated, the AIC or his/her designee (usually the Planning Section Chief) conduct an event Hot Wash, which is a brief facilitated session to obtain IMG members’ verbal feedback on the incident. This session should be conducted prior to the release of the majority of the IMG staff. This session is coordinated with the OEM Emergency Planning, Exercise, and Evaluation staff preparing the incident After Action Report (AAR) and feedback received provided for use in developing the report.

A final AAR draws conclusions from data collected. All FDA Centers and Offices should submit their input to the FDA OEM Director. All relevant parties within FDA receive the final report.
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C. Organization and Assignment of Responsibilities

This section identifies the general roles and responsibilities of FDA headquarters and field organizational components, including those that assist in preventing and protecting against, responding to, and recovering from all hazards. FDA staff work closely with one another and with governmental and industry partners to ensure the safety, efficacy, and security of FDA-regulated products that mitigate the public health effects of an emergency or disaster in the United States or worldwide, using novel and expeditious approaches to product regulation for optimized availability and use in all populations.

FDA functions are organized into four directorates:

1. The Office of Operations (OO) oversees administrative functions such as human resources (HR), facilities, information technology (IT), and finance.

2. The Office of Foods and Veterinary Medicine (OFVM) provides oversight of FDA’s food and feed programs as well as leads the implementation of the FDA Food Safety Modernization Act of 2011 (FSMA). OFVM includes CVM and the Center for Food Safety and Applied Nutrition (CFSAN) which includes the Coordinated Outbreak Response and Evaluation (CORE) Network.

3. The Office of Medical Products and Tobacco (OMPT) provides high-level coordination and leadership across the Centers for drugs, biologics, medical devices, and tobacco products. This Office also oversees special medical programs.

4. The Office of Global Regulatory Operations and Policy (OGROP) is focused on globalization and import safety of food and drug production and supply. The Office provides direction and support to ORA and the Office of International Programs (OIP).

The FDA product Centers are discrete management entities and are responsible for the regulation of a defined set of products. Their professional staff includes clinical, scientific, and regulatory experts. This expertise (analytical, laboratory, sampling procedures, subject matter expertise, and industry knowledge) is available for critical consultation should an emergency or disaster occur. The Centers are responsible for scientific evaluations, decisions in conjunction with ORA in response-focus based on scientific evaluations, and policy decisions within their respective program areas. FDA product Centers participate in emergency operations when the agency response includes, or may include, regulatory activities or products under their jurisdiction.

Figure C-1 outlines FDA’s overall organizational structure and is followed by a brief description of the emergency functions performed by all responsible organizational components. Refer to individual Center/Office emergency plans and procedural documentation, as applicable, for more detailed information on emergency actions.
Figure C-1. FDA Organizational Chart
C.1 OFFICE OF THE COMMISSIONER

The OC (see Figure C-2) provides centralized direction and management services for agency-wide programs to ensure that FDA’s consumer protection efforts are effectively managed within its legal and regulatory framework and those available resources are put to the most efficient use. The OC also provides policymaking, program direction, coordination and liaison, and expert advice to agency leadership and programs in support of FDA’s science-based work. The Office includes the Commissioner and Deputy Commissioners.

Office of the Chief of Staff

The Office of the Chief of Staff (OCoS) coordinates staff activities in the OC and serves as the principal liaison to HHS. The Office advises and provides integrated policy analysis and strategic consultation to the Commissioner and Deputy Commissioners, and other senior FDA officials, on activities and issues that affect significant agency programs, projects, and initiatives. OCoS provides senior-level leadership and guidance on issues and actions tied to the agency’s communications with HHS and the White House, including correspondence for secretarial signatures.

C.1.1 Office of the Executive Secretariat

The Office of the Executive Secretariat (OES) provides direct support to the Commissioner, including preparation of briefing material and background information for meetings, correspondence management and control, and preparation and transmittal of information advisories and alerts to HHS. The Office develops and maintains information to monitor the Commissioner’s and FDA’s goals and priorities.
In addition, OES is responsible for:

- **Dockets Management.** The Dockets Management staff will continue to post critical public information to the Federal Register in accordance with Federal law.

- **Freedom of Information Act (FOIA).** The Division of Freedom of Information will continue to respond to new requests and denials, provide appeals and litigation support and FOIA and Privacy Act policy, and will assist the Office of Media Affairs (OMA) as needed to review possible public announcements prior to their dissemination.

### C.1.2 Office of the Chief Counsel

The Office of the Chief Counsel (OCC) is composed of litigators and counselors. Litigators handle both civil and criminal cases, participating in case development; drafting pleadings, motions, and briefs; and conducting discovery and trials. Counselors provide legal opinions to the major programs of the agency—drugs, foods, biologics, devices, veterinary products, tobacco products, and enforcement. They participate in rulemaking proceedings, legislative matters, policy deliberations, and international negotiations. In addition, FDA attorneys are involved in explaining agency programs to Congress, regulated industry, and the public. In an emergency or disaster, OCC is responsible for, but not limited to, the following activities:

- Providing FDA organizational components with legal counsel on incident-related regulatory activities; examples include EUAs, marketing of medical countermeasures, and human subject protection (HSP).

- Providing FDA organizational components with legal counsel related to sharing of information with other government agencies and the public with proper protections (e.g., for trade secrets and confidential commercial information).

- Advising ORA and other relevant agency organizational components on inspections and enforcement matters related to foods, drugs, biologics, medical devices, radiological products, animal drugs/feed, cosmetics, and tobacco products.

- Coordinating with the HHS Office of the General Counsel (OGC) on FDA-related issues as appropriate (e.g., on grants and contracts and on legal issues arising from activities of other HHS Operating Divisions [OPDIVs] in relation to FDA activities).

- Working with other government agencies and their attorneys, Congress, and the U.S. Courts on FDA-related issues (e.g., the U.S. Department of Justice [DOJ], FBI, Federal Trade Commission [FTC], USDA, U.S. Environmental Protection Agency [EPA], U.S. Department of State [DOS], and the U.S. Postal Service [USPS]; State Attorney Generals’ offices; congressional oversight and appropriations committees; and Federal and State courts).

- Providing legal advice and assistance to the Office of the Secretary on matters within the expertise of the OCC.

### C.1.3 Office of Legislation

The Office of Legislation (OL) directs and manages FDA’s legislative needs, pending legislation, oversight activities, and congressional relations consistent with the mission of the agency. In an emergency or disaster, OL is responsible for, but not limited to, the following activities:

- Keeping Congress apprised of FDA actions in coordination with the HHS Office of the Assistant Secretary for Legislation.

- Responding to congressional Requests for Information (RFIs).
• Arranging and supporting congressional meetings, briefings, and hearings.

C.1.4 Office of Policy and Planning

The Office of Policy and Planning (OPP) provides advice to the Commissioner and other key FDA officials on matters relating to strategic direction, policy, development of regulations and guidance, legislative issues, planning and evaluation activities, and budget. OPP manages FDA’s Advisory Committee program, coordinates the publication of agency rules and notices in the Federal Register, serves as the agency focal point for policy development, and helps ensure agency components adhere to FDA policies and regulations. This Office participates with the Commissioner in the formulation of the budget, basic policies, and operational philosophy that guide FDA in effectively achieving its goals and meeting its responsibilities.

C.1.4.1 Office of Policy

The Office of Policy is responsible for advising the Commissioner and other key agency officials on matters relating to agency policy and on regulations and industry guidance development.

C.1.4.2 Office of Planning

The Office of Planning develops programs and systems to evaluate overall FDA program accomplishments against objectives and priorities, recommending changes as necessary.

C.1.5 Office of External Affairs

The Office of External Affairs (OEA) is responsible for the following activities:

• Serves as the central point of communication and education about FDA’s public health and regulatory activity. This includes the development, coordination, and leadership of all FDA communications and outreach efforts to the news media, health professionals, patient advocates, industry, States, consumer groups, and the general public. OEA also serves as the focal point for internal employee communications, speechwriting, creative and editorial services, and best practices in digital and web technology.

• Advises the Commissioner, Deputy Commissioners, and other key agency officials on FDA’s communications to the media, Congress, and the general public on issues that affect agency-wide programs, projects, strategies, partnerships, and initiatives.

• Advises and assists the Commissioner and other key officials on all public information programs; acts as the focal point for disseminating news on FDA activities and as a liaison with the U.S. Public Health Service (USPHS) and HHS on public information programs.

• Advises the Commissioner, Deputy Commissioners, and other senior staff throughout FDA on sensitive and controversial programs and initiatives that affect external stakeholder groups; provides historical expertise and records to inform those decisions.

• Provides communication expertise and state-of-the-art digital guidance and tools for application across the agency.

• Serves as a liaison between FDA and health professional and patient advocacy organizations to solve problems and address concerns these groups have with agency policies and programs related to human and medical product development and safety.

C.1.5.1 Web and Digital Media Staff

The Web and Digital Media Staff support the agency during an emergency with the following activities:
• Responsible for directing the design, content management, usability, and evaluation of the FDA website (www.fda.gov). Develops and interprets the agency’s web policies and serves as advocates for FDA’s web presence and catalysts for creative use of the web and digital media by the agency.

• Works closely, as partners, with the FDA Office of Information Management and Technology (OIMT), which is responsible for the technical operations of FDA’s website.

• Serves as the focal point and contact with the agency, HHS, and other Federal government website programs and operations.

• Provides direction, strategic planning assistance, and management coordination on agency website and digital media programs.

• Works closely with the website contacts in each of the Centers and principal Offices of the Commissioner to plan, coordinate, execute, and evaluate the agency’s website operations.

• Designs, develops, implements, monitors, and manages information published on the agency’s website and external digital assets.

• Delivers the agency’s messages to the public via the agency’s website and strategic online partnerships in the government, private, and nonprofit sectors.

• Establishes, manages, and monitors the implementation of agency standards for social media. Provides direction and strategic planning assistance related to social media.

C.1.5.2 Office of Communications

The Office of Communications supports the agency during an emergency with the following activities:

• Oversees and directs the agency’s print and online communications and visual identity to ensure quality and consistency as well as coherence in decision-making and the efficient operation of these internal functions across the agency and Department as a whole.

• Acts as the agency’s liaison with HHS for all publications and audiovisual needs; provides pre-publication clearance of publications, exhibits, and audiovisual materials in accordance with procedures established by the agency, USPHS, HHS, the Office of Management and Budget (OMB), and the White House.

• Creates and disseminates the agency’s flagship consumer health information, which includes timely and easy-to-read consumer update articles, videos, and photo slideshows containing the latest on all FDA-regulated products and practical wellness and prevention information to empower consumers. Manages the “For Consumers” section of the FDA website.

• Directs all aspects of FDA’s internal communications for employees, including the content and visual strategies for the homepage of the agency’s internal website; creates and coordinates agency-wide creation of content for reaching employees using a broad range of communications vehicles.

• Drafts speeches, informal remarks, talking points, slides, op-ed pieces, and letters to the editor for the Commissioner. Advises the Commissioner and other senior FDA executives on messages about FDA’s public health priorities, initiatives, and crucial priorities.

• Manages speaker requests for issues that cut across FDA’s organizational and product lines, as well as major meetings that involve various FDA Centers and Offices.
• Serves as a key resource for historical perspectives as well as records and resources used for agency communications and programs, including printed and online information, commemoratives, anniversaries, and milestones.

**C.1.5.3 Consumer Health and Constituent Affairs**

Consumer Health and Constituents Affairs support the agency during an emergency with the following activities:

• Advises the Commissioner and other key FDA officials on matters related to patients, patient advocacy, health professionals, consumers, State and Federal activities, and industry issues.

• Assists in the planning, administration, development, and evaluation of FDA policies related to patient advocacy and health professional organizations, consumers, States, and industry on serious and life-threatening issues.

• Serves as a liaison between FDA and stakeholder organizations to educate various constituents on FDA-related issues and activities.

• Provides internal coordination on FDA activities related to patient advocacy, health professional organizations, consumer groups, State organizations, and industry groups on high-priority topics such as serious and life-threatening diseases, imminent public health needs, and other special health issues.

• Coordinates and implements policies, programs, and initiatives related to MedWatch, including the MedWatch web pages, E-list, RSS [Really Simple Syndication] feed, and Twitter account.

• Ensures patient perspectives are taken into consideration during drug development and policy issues through the patient representative and patient consultant programs. Encourages and supports active participation of these groups in forming FDA regulatory policy to ensure the agency’s decisions are based upon a full range of perspectives.

• Administers MedWatch, the FDA safety information and adverse event reporting program. Also provides education and information on FDA’s key responsibilities, particularly the drug approval process, clinical trial design, and expanded access to investigational therapeutic products.

**C.1.5.4 Office of Media Affairs**

**C.1.5.4.1 Media Relations Staff**

The Media Relations Staff support the agency during an emergency with the following activities:

• Advises and assists the Commissioner of Food and Drugs and other key FDA officials on all news media activities; serves as a liaison with USPHS and HHS on news media activities.

• Serves as the agency focal point for preparing, clearing, and disseminating press announcements and other statements for the news media on agency activities, as well as arranging and facilitating press conferences, media briefings, media availabilities, interviews, and other news events.

• Establishes policy for and responds to news media inquiries with timely and accurate information; coordinates and maintains liaison with news media covering FDA activities.

• Facilitates news media interviews with senior agency officials and provides guidance and appropriate training to interview subjects on advisable conduct during news media contacts.

• Analyzes media coverage, conducts assessments and evaluations of media relations tactics to ensure objectives were met, and identifies areas for improvement.
• Plans, develops, and implements agency-wide multimedia communications strategies for disseminating regulatory and educational materials to the public through the news media in support of the agency’s top priorities and initiatives.

• Provides important daily newsclips about the agency to FDA employees and distributes the Daily News Media Report to HHS.

• Delegates FOIA denial authority to the Division of Freedom of Information for the agency.

C.1.6 Office of the Chief Scientist

The Office of the Chief Scientist (OCS) serves as the agency’s focus for scientific, medical, and related activities within the OC. OCS provides strategic leadership, coordination, and expertise to support scientific excellence, innovation, and capacity to achieve FDA’s public health mission. The Offices within OCS are as follows: the Office of Counterterrorism and Emerging Threats (OCET), Office of Critical Path Programs, and the newly established Office of Scientific Integrity (OSI) and Office of Regulatory Science and Innovation (ORSI). Additionally, the National Center for Toxicological Research (NCTR) has a direct reporting relationship to the OC and an indirect reporting relationship to OCS.

OCS also supports science and public health activities by effectively anticipating and responding to counterterrorism and emerging deliberate and natural threats (e.g., CBRN) to U.S. and global health and security, including through OCET.

C.1.6.1 Office of Counterterrorism and Emerging Threats

The Office of Counterterrorism and Emerging Threats (OCET) provides strategic leadership and coordination for FDA’s counterterrorism and emerging threat portfolios.

OCET performs the following main functions:

• Serves as FDA’s point of entry on policy and planning matters concerning global health security, counterterrorism and emerging threats

• Develops and coordinates the implementation of comprehensive FDA plans and strategies for these portfolios in collaboration with FDA Centers and Offices, and with external U.S. government and international partners

• Serves as the FDA focal point for the HHS Public Health Emergency Medical Countermeasures Enterprise disclaimer icon (PHEMCE) and Department of Defense medical countermeasure (MCM) programs to support the warfighter

• Coordinates FDA’s Medical Countermeasures Initiative (MCMi) to facilitate the development of safe and effective MCMs against chemical, biological, radiological, and nuclear agents and emerging threats, such as pandemic influenza

• Develops and coordinates implementation of FDA policies and procedures to facilitate the availability of MCMs, including efforts to safeguard MCMs from adulteration or disruption of supplies during public health emergencies, and enable access to available MCMs when necessary through an appropriate mechanism, such as Emergency Use Authorization

• On behalf of the Commissioner, facilitates communications within FDA and with external partners
C.1.6.2 National Center for Toxicological Research

NCTR conducts FDA mission-critical scientific research to support and anticipate FDA’s regulatory needs. This research is targeted to understand critical biological events in the expression of toxicity and to develop and characterize methods and incorporate new technologies to improve the assessment of human exposure, susceptibility, and risk. In an emergency or disaster, NCTR is responsible for, but not limited to, the following activities:

- Responding to RFIs from the Commissioner, other FDA Centers, and Federal agencies.
- Providing SME consultations to support informed decision-making in responding to health-related issues associated with the incident.

C.2 Office of Operations

OO provides executive direction, leadership, coordination, and guidance for the day-to-day operations of FDA; manages overall budgets and resources; and oversees management and business activities across all FDA headquarters and field offices. OO also ensures that proper conduct of FDA’s administrative and financial management activities, including budget, finance, acquisitions, IT, HR, organization, methods, and similar support activities effectively support program operations. These activities are available resources to support FDA’s emergency phases. The Offices under OO that report to the Chief Operating Officer and have responsibilities related to an emergency phase are:

- Office of Business Services (OBS)
- Office of Finance, Budget, and Acquisition (OFBA)
- Office of Human Resources (OHR)
- Office of Facilities Engineering and Mission Support Services (OFEMS)
- Office of Security and Emergency Management (OSEM)

In the Immediate Office, the Employee Safety and Environmental Management staff in an emergency or a disaster is responsible for, but not limited to, the following activities:

- Ensuring employees’ safety prior to, during, and while recovering from an emergency event by providing guidance and available resources to staff.
- Providing subject matter expertise and leadership for radiological emergencies.

C.2.1 Office of Business Services

OBS in an emergency or a disaster is responsible for, but not limited to, the following activity:

- Ensuring effective employee administrative support through Employee Resource and Information Center (ERIC) Call Center operations.

C.2.2 Office of Finance, Budget, and Acquisition

C.2.2.1 Office of Budget

The Office of Budget (OB) is responsible for developing budget formulation plans and organizes and carries out annual and multiyear budgeting in support of the nationwide public health protection programs administered by FDA. In coordination with FDA Centers and Offices, OB responds to requests for budget information, special reports, and exhibits and provides expertise and performs all duties involved with the formulation, justification, and presentation of FDA budget submissions to HHS, OMB, and Congress. In an emergency or disaster, OB is responsible for, but not limited to, the following activities:
• Providing expert advice on and managing supplemental budget requests to support unforeseen expenses during critical periods or events; OB would facilitate this process by working with senior management and Centers to identify resource needs, implementing the process to formulate the budget request, providing essential support during the review and clearance process, and responding to questions from Congressional Appropriations staff and other stakeholders.

• Facilitating that process by conducting outreach and coordinated communications; gathering information; and communicating with senior FDA management, Centers, HHS, OMB, and Congressional Appropriations staff.

C.2.2.2 Office of Acquisitions and Grants Services

The Office of Acquisitions and Grants Services (OAGS) is responsible for negotiating, awarding, and managing all contracts, cooperative agreements, interagency agreements, grants, administration of the purchase card program for FDA, MOUs, and technology transfers. OAGS is also responsible for writing acquisition policy, providing strategic business advice and support to our stakeholders, and liaising with the HHS Senior Procurement Executive.

Administrative functions, such as payroll, travel, and timekeeping, will be provided by the Administrative Officer, Director’s Administrative Assistant, and other administrative support personnel.

C.2.2.3 Office of Financial Operations

The Offices that comprise the Office of Financial Operations (OFO) plan, direct, and coordinate a comprehensive financial management operations program for FDA, encompassing the areas of budget analysis, execution, automated financial systems, fiscal accounting, internal financial audit, financial services related to accounts payable, travel support and payroll liaison, and financial reporting.

C.2.2.3.1 Office of Financial Management

The Office of Financial Management (OFM) is FDA’s steward of financial assets and resources. OFM performs the following essential functions to support FDA’s response to an emergency:

• **Budget Execution.** Apportions funds appropriated by Congress among components and oversees transfers of funds between components; this includes processing user fees.

• **Accounts Receivable – User Fees.** Receives user fee payments, which allows FDA’s certification process to continue.

• **Financial Statements and Reports.** Prepares the agency’s financial statements and maintains its general ledger.

C.2.2.3.2 Office of Financial Services

The Office of Financial Services is responsible for the following activities:

• Directs and coordinates operations for financial services related to accounts payable, travel support, and payroll liaison.

• Monthly, Quarterly, and Yearly Closing—coordinates month-end, quarter-end, and year-end close of financial operations within the Unified Financial Management System (UFMS).

• Maintains liaison with the Program Support Center (PSC) and the Defense Financial Accounting System (DFAS) representatives on issues relating to pay and leave and monitors the processes to ensure the successful payment to employees.
• When necessary during an event, the Division of Travel Services may oversee the following: processing of vouchers and traveler’s reimbursements, the Concur Government Edition (CGE) system, the agency’s Travel Card and Centrally Billed Account programs, and travel process for all State employees working in tandem with ORA employees.

### C.2.3 Office of Information Management and Technology

OIMT enables FDA’s strategic efforts to transform and improve the information systems and infrastructure needed to support critical agency operations. The Office implements and enhances common systems to support FDA’s regulated products. It also maximizes the availability and use of information technologies that increase or enhance electronic access for the public, as well as the full span of FDA’s external and internal customer base, while maintaining effective security. OIMT aligns IT investments to business goals that fully support core mission and business priorities and reduces costs of existing legacy systems. The Office consolidates, modernizes, and optimizes FDA’s IT infrastructure and provides the platform required for the agency to meet IT initiatives and to move towards the bioinformatics era of science-based decisions.

In an emergency or disaster, OIMT is responsible for, but not limited to, the following activities:

- Ensuring the command and control of all reporting to OIMT.
- Ensuring data centers (there are four main data centers) are functioning and coordinating with the appropriate building management to determine the building status.
- Ensuring IT security services are available that are essential to protect FDA’s IT assets and identifying and applying patches that will address the most critical security vulnerabilities.
- Ensuring the availability of FDA local area network (LAN)/wide area network (WAN) that will be critical for both field and local users to access agency IT resources.
- Maintaining Internet services that are critical to the agency’s ability to communicate with regulated industry and providing access to FDA’s internal and external IT resources.
- Ensuring Active Directory availability to provide authentication services when users log onto FDA systems.
- Ensuring telephone availability, Voice over Internet Protocol (VoIP) customer support, and continuity of agency email/BlackBerry services.
- Monitoring the availability of agency Outlook Web Access (OWA).
- Managing and maintaining IT help desk and application support services.
- Ensuring a Special Routing Arrangement Service infrastructure that will support 16,000 concurrent broadband users and approximately 350 dial-in users.
- Ensuring FDA staff can perform non-standard queries and data extractions.
- Ensuring agency computer application and file/print services.
- Monitoring and maintaining the availability of applications and databases.
- Procuring additional contract services or information system hardware/software components as appropriate.

### C.2.4 Office of Human Resources

In an emergency or disaster, OHR is responsible for, but not limited to, the following activities:
• Reviewing employee issues related to telework, leave, flexiplac, social distancing, and other HR issues; assessing the impacts of related decisions, such as building closures; issuing clear guidance consistent with the Office of Personnel Management (OPM), HHS, and other administration directives and policies while integrating program priorities and mission requirements; and maintaining a clear line of communication on both an agency and individual level so that the implementation of policies is consistent and well understood across FDA.

• Coordinating and liaising with the FDA Office of Financial Services to ensure OHR employees adhere to any timekeeping updates necessary to continue receiving compensation.

• Providing coordination between FDA management and the Assistant Secretary for Health’s Commissioned Corps programs; serving the FDA Centers, special assignments, and details to other organizations and initiatives.

• Providing leadership and direction regarding all aspects of agency-wide HR management.

• Ensuring internal and external customer services continue to be provided.

C.2.5 Office of Facilities Engineering and Mission Support Services

In an emergency or disaster, OFEMS is responsible for, but not limited to, the following activities:

• Ensuring FDA facilities are safe and available for staff and acquiring temporary facilities if necessary.

• Providing direct interface with the U.S. General Services Administration (GSA) for services in all leased facilities.

C.2.6 Office of Security and Emergency Management

OSEM is responsible to assure that FDA is supported in the areas of personnel and physical security, and emergency management through a coordinated effort among these functional areas. This includes planning for continued operations in the face of a crisis and assuring that public health is protected to the greatest extent possible through coordinating emergency response activities involving FDA-regulated products. OSEM also coordinates agency COOP activities, which address how to continue agency operations when the availability of FDA facilities to accomplish agency work has been impacted by an emergency.

C.2.6.1 Office of Security Operations

OSO has responsibility for agency-wide physical and personnel security programs, including the suitability and National Security Information program. In an emergency or disaster, OSO is responsible for, but not limited to, the following activities:

• Ensuring the national security clearances of personnel assigned to various EOCs as part of a response are either sent to or received by the appropriate EOC.

• Providing guidance on the proper handling, marking, processing, and storage of classified materials.

• Ensuring the physical security of all nationwide FDA facilities; the Physical Security Branch may coordinate with on-duty security guard personnel regarding any temporary staff deployment issues that may arise during an emergency.
C.2.6.2   Office of Emergency Management

OEM serves as FDA’s focal point for coordinating emergency response activities involving FDA-regulated products and in situations when agency resources need to be used or deployed. It coordinates intra- and interagency activities related to emergency preparedness and response and security operations. OEM assists in the development, management, and coordination of incident management policies and programs for FDA to ensure that a structure exists to respond rapidly and effectively to all hazards. OEM consists of a Director; Immediate Office; Program Operations and Coordination Staff (POCS); Emergency Planning, Exercises, and Evaluation Staff (EPEES); and OEM/OEO. OEM/POCS provides GIS support to the agency and manages the FDA After-Hours Emergency Call Center, GeoWeb (a geographic information portal), and the Emergency Operations Network – Incident Management System (EON-IMS). OEM/EPEES is responsible for emergency plans, exercises, and evaluation. Additionally, OEM serves as a coordinator to the HHS Office of the Assistant Secretary for Preparedness and Response (ASPR), providing situational awareness of all FDA-related emergencies, and ensures FDA’s emergency operations procedures are in alignment with national and HHS procedures. During an emergency response, OEM supports the agency with the following activities:

- Coordination, monitoring, documentation, and reporting on all agency response activities and interagency communications through the Office of Emergency Operations (see Section C.2.6.2.1).

- Provides Offices and Centers with maps and spatial analysis through GIS for use in strategic planning of agency emergency response activities. Additionally, GIS provides situational awareness during a response in the form of maps that depict FDA-regulated industry, locations of FDA response activities, and other areas of interest. GIS allows responders to explore spatial trends in information and visually communicate information that can support and improve decisions.

- Oversees the FDA Emergency Call Center, which provides service for responding to public inquiries and reports related to FDA-regulated products as well as surge capacity service for managing increased volumes of inquiries due to an incident involving an FDA-regulated product.

- Manages FDA’s EOC through OEO (see Section C.2.6.2.1).

- Coordinates the agency’s evaluation or after action review of FDA’s responses to incidents and emergencies to determine appropriate internal and external referral for further action and recommended changes in agency policies and procedures.

- Advises and assists key FDA officials on counterterrorism and intelligence matters. Serves as lead for the agency’s Special Events planning and preparedness activities.

C.2.6.2.1   Office of Emergency Operations

OEO is staffed with emergency coordinators and serves as the central headquarters coordination point for the agency’s response to adverse events and injuries associated with regulated products, foodborne illnesses, product tampering, and manmade and natural hazards. It serves as the agency focal point for emergency preparedness and response operating the 24/7 emergency response system, and manages the FDA Emergency Operations Center (EOC). OEO reviews and assists in the analysis of preliminary information about threats and hazards, and assists in the early recognition and agency notification of emergencies, outbreaks, natural disasters, and terrorism or other criminal acts. Part of OEO responsibilities is to manage the national Consumer Complaint System, which monitors reports of problems with FDA-regulated products for potential emergencies. OEO also works across the agency at the Headquarters level for planning and execution of FDA’s activities in Special Events, including National Special Security Events (NSSSEs). Another responsibility is to participate in weekly National Biosurveillance Integration Center (NBIC) conference calls sponsored by DHS to provide a secure forum
for interagency information sharing for early recognition of biological events of national concern, both natural and manmade, to make a timely response possible.

In an emergency or disaster, OEO’s staff members are responsible for, but not limited to, the following activities:

- Coordinating, monitoring, documenting, and reporting on all agency response activities and inter-agency communications.
- Coordinating resource requirements with headquarters Centers and Offices and issuing assignments to the field.
- Managing FDA’s EOC and staffing it with personnel from OEM/OEO as operational status requires. When the EOC is activated, OEM may augment EOC operations with personnel from relevant Centers and Offices to monitor emergency situations, triage complaints and alerts, issue MAs to organizational components, coordinate overall agency response operations, and communicate with external partners requesting technical and material support.
- Maintaining situational awareness on intelligence matters that may have an impact on FDA staff, facilities, and/or regulated products.
- Triaging consumer complaints and alerts.
- Collecting and disseminating situational awareness information on natural and manmade disasters.
- Communicating with Federal agencies (e.g., HHS ASPR and the HHS Secretary’s Operations Center, CDC, USDA, DHS, National Oceanic and Atmospheric Administration [NOAA], EPA) as they request technical and material support from FDA.
- Providing essential staff for the operation of FDA’s EOC and liaisons to the HHS SOC.

C.3 Office of Foods and Veterinary Medicine

The FDA Foods and Veterinary Medicine (FVM) Program includes CFSAN and CVM, along with the food-related activities of ORA and the CORE Network, and draws on the resources and expertise of FDA’s NCTR and key OC staff Offices.

OFVM provides executive leadership and strategic direction to the FVM Programs to protect and promote the health of humans and animals by ensuring the safety of the American food supply, food additives, and dietary supplements as well as the safety of animal feed and the safety and effectiveness of animal drugs. The FVM Program does this by setting science-based standards for preventing food and feedborne illnesses and ensuring compliance with these standards, protecting the food and feed supply from intentional contamination, and ensuring that food labels contain reliable information and encouraging product reformulation to allow consumers to make healthy choices and promote well-being. OFVM is also responsible for leading FDA’s efforts in responding to foodborne outbreaks and contamination. The FVM Program leads the implementation of the FDA FSMA. The FVM Program also promotes and protects the health of humans and animals by regulating the manufacture and distribution of food additives and drugs that will be given to animals.

The FDA FVM Program protects and promotes the health of humans and animals by:

- Posting vital and important information, emergency messages, and other communication information quickly to users via the FDA website, in coordination with OEA.
- Ensuring the safety of foods for humans, including dietary supplements.
- Ensuring the safety of animal feed and the safety and effectiveness of animal drugs.
Setting science-based standards for preventing foodborne illness and ensuring compliance with these standards.

Coordinating foodborne outbreak surveillance, response, and post-response activities related to incidents involving multiple illnesses linked to FDA-regulated human and animal food and cosmetic products.

Protecting the food and feed supply from intentional contamination.

Ensuring that food labels contain reliable information consumers can use to choose a healthy diet.

### C.3.1 Center for Food Safety and Applied Nutrition

CFSAN, in conjunction with the agency’s field staff, is responsible for promoting and protecting the public’s health by ensuring that the Nation’s food supply is safe, sanitary, wholesome, and honestly labeled and that cosmetic products are safe and labeled properly. CFSAN has the authority to regulate establishments that manufacture, process, pack, hold, or grow food involved in interstate commerce, including manufacturers, distributors, and warehouses. It also determines whether data collected by another agency or organization are adequate for FDA decisions regarding food and cosmetic issues. In an emergency or disaster, CFSAN is responsible for, but not limited to, the following activities:

- Collaborating with other public health agencies and industry regarding food contaminants and other monitoring programs for foodborne illness programs.
- Providing critical information on food safety, food defense, and regulatory issues to consumers, industry, and other Federal and SLTT governmental and international entities (all levels of the food chain).
- Protecting human health through regulatory, legal, and administrative actions.
- Participating actively in legal proceedings critical to the safety and defense of the food supply.
- Using import alerts to prevent unsafe food and cosmetic products from entering the United States.
- Facilitating coordination of food contamination investigations between FDA and the States.
- Identifying foods that are at elevated risk of contamination and investigating the effectiveness of food processing and preparation practices.
- Developing and disseminating recommendations on measures to prevent the contamination of FDA-regulated foods and cosmetics.
- Developing and evaluating analytical methods for identifying food and cosmetic contaminants.
- Critically reviewing cases and other regulatory actions sent from the Office of Compliance to evaluate whether proposed regulatory action is supported by findings and providing policy and technical input to FDA field investigations.
- Identifying and training a cadre of CFSAN personnel for potential rapid deployment to the FDA EOC, to HHS, and to external organizations to provide support for contingency functions.

### C.3.1.1 Coordinated Outbreak Response and Evaluation Network

The CORE Network is responsible for coordinating foodborne outbreak surveillance, response, and post-response activities related to incidents involving multiple illnesses linked to FDA-regulated human food and cosmetic products. Members of the CORE Network at the FDA headquarters level are assigned on a continuous, full-time basis to multidisciplinary teams that coordinate efforts internally within FDA, including CFSAN, CVM, ORA, and OFVM, as well as with external stakeholders, including the CDC,
USDA, and State and local health, agriculture, and regulatory agencies. The network is led by the OFVM Chief Medical Officer (CMO)/Director of CORE, who has the overall responsibility for leadership and management of FDA’s activities related to incidents of illness linked to human food and serves as the spokesperson during outbreaks.

CORE Network leadership includes a Deputy Director, who participates fully with the CMO in providing the leadership, policy development, decision-making, and strategic planning for food-related outbreaks and food safety issues affecting the public health within the purview of FDA. Additionally, two managers oversee the areas of Prevention (includes the Signals and Surveillance and Post-Response) and three Response Teams. Also included in the network are key strategic FDA resources in both the field and at FDA headquarters (the District Offices, the Rapid Response Teams [RRTs] that work with State partners; FDA’s OEA; the SMEs at CFSAN and CVM; and others as appropriate). In specific instances, the CORE Network will work with OEM to determine whether an IMG should be activated if the response capacity of CORE is exceeded.

In specific incidents, OEM/OEO will be immediately notified if the size, severity, and/or scope of the response to an incident is above the capacity of CORE. In this case, CORE will request the assistance of the OEM to activate an IMG to coordinate the response to the incident consistent with the FDA EOP.

C.3.2 Center for Veterinary Medicine

CVM ensures the safety, efficacy, and quality of drugs for animals (including food-producing and companion animals), animal food and feed, and medical devices used on animals potentially threatened by an incident. In an emergency or disaster, CVM is responsible for, but not limited to, the following activities:

- Protecting animal and human health through regulatory, legal, and administrative actions.
- Actively participating in legal proceedings critical to the safety of the food and feed supply.
- Using import alerts to prevent unsafe products from entering the United States.
- Facilitating coordination of feed, meat, and milk residue investigations between FDA and the States.
- Providing sponsors and other stakeholders with guidance on the use of animal drugs that serve as potential medical countermeasures in animals.
- Providing information on critical safety and regulatory issues to stakeholders, consumers, industry, veterinarians, and other government officials.
- Identifying feeds that are at elevated risk of contamination and investigating the effectiveness of feed processing and preparation practices.
- Coordinating human and animal foodborne outbreak surveillance, response, and post-response activities related to incidents involving FDA-regulated animal food.
- Coordinating animal outbreak surveillance, response, and post-response activities related to incidents involving FDA-regulated animal devices.
- Collaborating with public health agencies regarding feed contaminant, tissue residue programs, and other monitoring programs for meat and poultry products.
- Providing information regarding manufacturers’ compliance with Good Manufacturing Practices (GMPs) and other relevant animal drug product quality issues.
- Providing technical advice and assistance in the assessment of animal drug or feed product possibly affected.
• Assessing the availability, production capacity, and surge capacity of animal drugs and providing information on alternative sources of critical products in shortage situations.

• Providing an assessment on the diversion of contaminated human food for animal feed use.

• Providing advice to pet owners regarding animal safety measures.

C.4 Office of Medical Products and Tobacco

OMPT provides executive leadership, management, and policy direction to all FDA medical product- and tobacco-related programs. The Office exercises, on behalf of the Commissioner, direct line authority over CDER, CBER, CDRH, and the Center for Tobacco Products (CTP). The Office directs the activities of FDA’s special medical programs, including OCP, the Office of Orphan Products Development, the Office of Good Clinical Practice (OGCP), and the Office of Pediatric Therapeutics (OPT). The Office directs efforts to integrate the programs, as necessary, of CDER, CBER, CDRH, and CTP, and thereby ensure the optimal use of all available FDA resources and tools to improve the safety and proper labeling of medical products and tobacco. The Office is responsible for the following activities:

• Directs the development of integrated strategies, plans, policies, and budgets to build FDA’s medical product- and tobacco-related scientific and regulatory capacities and programs, including recruitment and training of key personnel and development of information systems.

• Represents FDA on medical product- and tobacco-related matters in dealing with the Office of the Secretary of HHS, the White House, other elements of the Executive Branch, and with Congress.

• In conjunction with the Deputy Commissioner for Global Regulatory Operations and Policy, represents FDA on medical product- and tobacco-related matters in dealing with foreign governments and international organizations.

• Directs FDA efforts to build consistency and uniformity in the evaluation for safety and efficacy of new medical products and new tobacco products.

C.4.1 Office of Special Medical Programs

The Office of Special Medical Programs (OSMP) Immediate Office oversees and coordinates the program offices that comprise OSMP. The OSMP Immediate Office:

• Provides leadership and direction in the coordination of internal and external review of pediatric science, safety, ethics, and international issues in accordance with laws and regulations.

• Oversees the implementation of the Orphan Products provisions of the FD&C Act to encourage the development of drugs of limited commercial value for use in rare diseases and conditions to promote the public health; the Immediate Office provides leadership in developing and communicating agency policy on orphan product activities, including protocol assistance, designation, exclusivity, and grants and contracts.

• Provides leadership and direction on Good Clinical Practice (GCP) and HSP regulation, policy, harmonization, and outreach activities; the Immediate Office oversees development of regulations and policy to help ensure protection of human subjects involved in FDA-regulated research and the integrity of data resulting from such trials.

• Provides leadership and direction on issues involving the regulation of combination products, the classification of human medical products, and jurisdiction over human medical products; the Immediate Office oversees development of regulations, policy, procedures, and processes to
facilitate classification of human medical products and the agency’s regulation, review, and oversight of combination products.

- Provides leadership in management and oversight of FDA Advisory Committees to provide consistent application of laws and policies applicable to such committees; the Immediate Office oversees development of policy, procedures, and processes to maintain and improve the agency’s Advisory Committee program.

- Evaluates and responds to appeals involving product classification decisions, assignment of combination product decisions, and orphan product designations.

C.4.1.1 Office of Good Clinical Practice

In general, the mission of the OGCP focuses on longer term policy/regulation development and intra-agency coordination of issues related to the conduct of clinical trials and the protection of the rights, safety, and welfare of subjects in FDA-regulated studies. There are, however, several mission-critical areas for which OGCP is responsible that require daily attention and that may have heightened importance during an emergency situation, particularly an emergency that directly impacts the conduct of ongoing clinical trials. These areas include addressing clinical trial/HSP inquiries, questions related to emergency or compassionate use of investigational articles, trial-specific concerns, EUAs, and coordination of interagency GCP/HSP issues. Should the emergency situation have a major impact on the conduct of clinical trials in one or more areas of the country, OGCP could potentially establish an emergency-specific hotline, as was done in the aftermath of Hurricane Katrina. Major concerns regarding subject safety could include identification of an accessible clinical site(s) for continued subject care depending on the specific situation.

C.4.1.2 Office of Combination Products

OCP ensures the prompt assignment of combination products to FDA Centers, the timely and effective pre-market review of such products, and consistent and appropriate post-market regulation of like products subject to the same statutory requirements to the extent permitted by law. In addition, OCP is responsible for determining whether articles are drugs, devices, biological products, or combination products if their classification is unclear or in dispute. In an emergency or disaster, OCP is responsible for, but not limited to, the following activities:

- Classifying articles as drugs, devices, biological products, or combination products.
- Assigning an FDA Center to have primary jurisdiction for review of a combination product.
- Ensuring consistency and appropriateness of post-market regulation of combination products.
- Updating agreements, guidance documents, or practices specific to the assignment of combination products during the incident.
- Working with FDA Centers to implement guidance or regulations to clarify the agency regulation of combination products.
- Serving as a focal point for combination products issues for internal and external stakeholders.

C.4.1.3 Office of Orphan Products Development

The Office of Orphan Products Development develops and communicates agency policy and makes decisions on approval of sponsor requests and incentives, including orphan product protocol assistance, orphan product designation, orphan product exclusivity, and orphan product grants and contracts, to support clinical research and other areas of agency policy related to the development of products for rare disorders. It reviews IND and biologics applications and IDEs to locate the existence of products under investigational study that show promise for effectiveness for rare or common diseases but lack
commercial sponsorship. It assists sponsors, researchers, and investigators in communicating with agency regulatory officials and expediting solutions to problems in obtaining investigational exemptions or pre-market approval status.

C.4.1.4 Office of Pediatric Therapeutics

OPT coordinates and facilitates all activities of FDA that may have any affect on a pediatric population or the practice of pediatrics or may in any other way involve pediatric issues. In an emergency or disaster, OPT is responsible for, but not limited to, the following activities:

- In direct coordination with CDER’s Office of Surveillance and Epidemiology and CDER’s Pediatric and Maternal staff (within the Office of New Drugs), and in consultation with CDER’s Office of Counterterrorism and Emergency Coordination, review and address all adverse event reporting involving drugs used in the pediatric population.
- In direct coordination with CDRH’s Pediatric Steering Committee and Medical Product Surveillance Network (MedSun), address emergency issues related to device use in pediatrics.
- Provide consultation and coordination on pediatric issues across Centers and with external groups and agencies as well as provide representation on groups such as the Public Health Emergency Medical Countermeasures Enterprise (PHEMCE) working group.

C.4.1.5 Advisory Committee Oversight and Management Staff

The Advisory Committee Oversight and Management Staff (ACOMS) coordinates and facilitates all activities of FDA that are related to the establishment and operations of FDA’s Advisory Committees. ACOMS is responsible for seeing that all statutory and regulatory requirements and agency policies are met when conducting all Advisory Committee meetings, including the screening of members for potential financial conflicts of interests. In an emergency or disaster, ACOMS is responsible for the following activities, among others:

- Overseeing the operations of the Centers to obtain the appropriate experts to attend a meeting.
- Assisting the Centers, if necessary, in procuring a space to hold the meeting.
- Assisting the Centers, the Regulations Policy Management staff, and the Associate Commissioner for Special Medical Programs in publishing public meeting announcements in the Federal Register and on the FDA website.
- Reviewing the financial interests of all individuals invited to participate in the meeting for potential financial conflicts of interest. (This is conducted with the assistance of the Centers, the Ethics and Integrity staff, and the Associate Commissioner for Special Medical Programs. If any members require waivers to participate, ACOMS coordinates disclosure with the Division of Freedom of Information and the FDA web staff.)

C.4.2 Center for Drug Evaluation and Research

CDER ensures the safety, efficacy, and quality of prescription and over-the-counter drugs, (both brand name and generic), and therapeutic biological agents. In an emergency or disaster, CDER is responsible for, but not limited to, the following activities:

- Providing sponsors and other stakeholders with regulatory guidance on the development and use of medical countermeasures against CBRNE agents or emerging infectious threats (e.g., severe acute respiratory syndrome [SARS] and 2009 H1N1 [swine flu]) when there is no FDA approval for the product or proposed indication or when another issue would cause the product to be considered adulterated or misbranded.
- Facilitating communications with manufacturers on adverse event reporting during an emergency.
- Collaborating with public health agencies regarding drug and therapeutic biologic product stockpile issues, including labeling, appropriate usage, product performance, monitoring, use in special populations, product expiry (e.g., the Shelf Life Extension Program), and other evolving issues.
- Processing EUA requests for drugs and therapeutic biologic agents after a public health emergency is declared by the Secretary.
- Providing information to manufacturers regarding compliance with GMP and other relevant product quality issues, including extension of product expiry.
- Assessing the availability, production capacity, and surge capacity of drugs and therapeutic biologic products used as countermeasures and providing information on alternative sources of critical medical countermeasures in shortage situations.
- Managing CDER laboratory capabilities.
- Monitoring media and marketplace for counterfeit or violative drugs (or advertising) of relevance to the emergency.
- Enhanced monitoring and evaluation of adverse events associated with products relevant to the emergency.
- Reviewing public communications related to CDER-regulated products relevant to the emergency.
- Investigating potential root causes of adverse events or product quality issues reported with drug and therapeutic biologic products.

C.4.3 Center for Biologics Evaluation and Research

CBER ensures the safety, efficacy, and quality of biological products (including blood and blood products, vaccines, tissues, and cellular and gene therapies) potentially used as medical countermeasures. In an emergency or disaster, CBER is responsible for, but not limited to, the following activities:

- Providing sponsors and other stakeholders with guidance on the use of biological products as medical countermeasures.
- Using sound science and regulatory expertise to review, evaluate, and license new biological products that are safe, effective, and of high-quality standards.
- Facilitating the availability of safe and effective vaccines, blood and blood products, cells, tissues, gene-therapies, devices, and other biological products used to prevent and treat disease outbreaks.
- Collaborating with public health agencies regarding biologics stockpiling issues, including labeling, appropriate usage, product performance, monitoring, use in special populations, and other evolving issues.
- Ensuring compliance with current GMP for all regulated manufacturers.
- Providing regulatory guidance on the use of unapproved products or unapproved uses of approved products.
- Processing EUA requests for biological products after a public health emergency is declared by the Secretary.
• Providing information regarding manufacturers’ compliance with current GMP and other relevant product quality issues.

• Assisting in the assessment of potentially contaminated biological products.

• Assessing the availability, production capacity, and surge capacity for biologic products used as medical countermeasures and providing information on alternative sources of critical medical countermeasures in shortage situations.

• Managing CBER laboratory testing and lot release capabilities.

• Communicating information, including risk information, about biological products to industry, consumers, SLTT governments, and other stakeholders.

• Conducting post-marketing surveillance and enforcement activities relating to biological products.

C.4.4 Center for Devices and Radiological Health

CDRH ensures the safety, efficacy, and quality of medical devices, including IVD and radiological products. In an emergency or disaster, CDRH is responsible for, but not limited to, the following activities:

• Processing EUA requests for medical devices (i.e., diagnostics, personal protective equipment [PPE], and radiological products) after a public health emergency is declared by the Secretary.

• Facilitating the development, regulatory review, and production of diagnostic devices, PPE, and other devices that may be needed.

• Conducting post-market surveillance to monitor the safety and effectiveness of devices used, including for diagnosis, therapeutics, PPE, and supportive care.

• Supporting efforts to ensure an adequate supply of devices through cooperative interactions with manufacturers and distributors and coordination with the Strategic National Stockpile (SNS) to determine the adequacy of stocks and actions required to meet targeted amounts.

• Using an emergency shortages database to identify and monitor supplies of certain devices that are or have the potential to be in demand.

• Educating device users on how to use devices in a safe and effective manner.

• Collaborating with public health agencies regarding product stockpile issues, appropriate usage, labeling, product performance, monitoring, use in special populations, and other evolving issues.

• Communicating information, including risk information, about medical devices and radiation-emitting products to industry, consumers, SLTT governments, and other stakeholders.

C.4.5 Center for Tobacco Products

CTP is responsible for protecting the public’s health by restricting the marketing and access of tobacco products to minors; regulating the manufacturing, distribution, and labeling of tobacco products; setting tobacco products standards; requiring registration of tobacco manufacturers; and requiring ingredient and constituent listings for tobacco products. In an emergency or disaster, CTP is responsible for, but not limited to, the following activities:

• Collaborating with other public health agencies to communicate information regarding tobacco product contaminants and to coordinate monitoring for patterns of exposure and illness.

• Protecting human health through regulatory, legal, and administrative actions.
• Participating actively in legal proceedings critical to ensuring compliance with tobacco product regulatory standards.
• Facilitating coordination of tobacco product contamination investigations between FDA and SLTT governmental and international entities.
• Identifying tobacco products that are at elevated risk of contamination and investigating the effectiveness of tobacco processing and preparation practices.
• Developing and disseminating recommendations on measures to prevent the contamination of tobacco products.
• Developing and evaluating analytical methods for identifying tobacco product contaminants.
• Critically reviewing cases and other regulatory actions sent from the Office of Compliance to evaluate whether proposed regulatory action is supported by findings and providing policy and technical input to FDA field investigations.
• Enhanced monitoring and evaluation of adverse events associated with products relevant to the emergency.
• Communicating information, including risk information, about tobacco products to industry, consumers, SLTT governments, and other stakeholders.
• Ordering a stop to the distribution of a tobacco product and recalling a tobacco product relevant to the emergency.

C.5 Office of Global Regulatory Operations and Policy

OGROP provides executive oversight, strategic leadership, and policy direction to FDA’s domestic and international product quality and safety efforts, including global collaboration, global data-sharing, development and harmonization of standards, field operations, compliance, and enforcement activities. GROP is responsible for the following activities:

• Drives improvement in FDA’s risk analytics and IT capability, in conjunction with OIMT, to enhance FDA’s gathering and use of intelligence in directing its regulatory efforts.
• Leverages public and private sector third parties to increase FDA’s global knowledge and coverage.
• Convenes and chairs a Global Programs Board consisting of the ACRA, the Directors of all FDA product Centers, and other agency leaders to coordinate and enhance FDA’s global efforts.
• Provides executive leadership and management to all programs in ORA and OIP.
• Exercises, on behalf of the Commissioner, direct line authority over ORA and OIP.
• Exercises, on behalf of the Commissioner, all international and ORA-related legal authorities that the Commissioner is empowered to exercise under the FD&C Act, as amended; the PHSA; and other applicable laws.
• Represents FDA on matters relating to domestic and international product quality and safety efforts in dealings with the Executive Branch, Congress, foreign governments and international organizations, and others.

C.5.1 Office of International Programs

OIP serves as the agency focal point for international issues. The Office leads, manages, and coordinates all of FDA’s international activities to effect an affirmative and strategic public health agenda in the
international area; enhance and maximize FDA’s communications and interactions globally, ensuring they reflect the agency’s policies and best scientific, legal, and policy thinking; ensure that FDA international communications and interaction are consistent with HHS public health objectives; and leverage resources with counterpart foreign agencies and international organizations in meeting FDA’s public health mission. In an emergency or disaster, OIP is responsible for, but not limited to, the following activities:

- Managing foreign offices and FDA staff stationed overseas.
- Coordinating FDA communications with DOS, including U.S. embassies abroad, foreign governments, and international organizations, including foreign embassies and consulates in the United States.
- Exchanging information, and advising on clearing such, with foreign counterparts and other international partners to ensure compliance with laws and policies and consistency in responses.
- Assessing in real-time and in-country, where possible, the conditions and events in those areas that might have an impact on the safety, quality, and availability of FDA-regulated products exported to the United States.
- Engaging more proactively and consistently with various international communities in strategic regions abroad, including foreign governments, foreign regulatory counterparts, international organizations, other U.S. government colleagues working abroad, industry, and the academia/research community; such engagements are intended to help FDA better accomplish its mission to collaborate with foreign regulatory authorities to reduce regulatory burdens, harmonize regulatory requirements, and establish appropriate reciprocal arrangements in order to promote and protect the public health of the United States by obtaining better information to help FDA Centers, ORA, and OC make better informed regulatory and other decisions.
- Managing formal arrangements with foreign governments (e.g., MOUs, mutual recognition agreements, exchange of letters, confidentiality commitments) to facilitate rapid and efficient information exchange and other cooperation.
- Facilitating international technical cooperation and assistance activities and capacity building for emergency collaborations.
- Implementing policies and procedures pertaining to emergency international travel and processing international travel requests.
- Facilitating international visitors as needed.
C.5.2 Office of Regulatory Affairs

ORA protects consumers and enhances public health by maximizing compliance of FDA-regulated products and minimizing risk associated with those products. ORA is led by the ACRA and is the lead Office for all agency field activities. ORA is composed of: Office of Management; Office of Communications and Project Management; Office of Criminal Investigation; Office of Training Education, and Development; Office of Partnership and Operational Policy; Office of Human and Animal Food Operations; Office of Medical Products and Tobacco Operations; Office of Enforcement and Import Operations; and Office of Regulatory Science. In an emergency or disaster, ORA’s responsibilities include, but are not limited to, the following activities:

- Coordinating, interpreting, and evaluating the agency’s overall compliance efforts.
- Providing advice or assistance to the Commissioner and other key officials on regulations and compliance-oriented matters related to the incident.
- Stimulating awareness within the agency of the need for prompt and positive action to ensure compliance by regulated industries.
- Working to ensure an effective and uniform balance between voluntary and regulatory compliance and agency responsiveness to consumer needs.
- Evaluating and coordinating all proposed legal actions to ascertain compliance with incident-related regulatory policy and enforcement objectives.
- Executing direct line authority over all agency field operations; developing, issuing, approving, or clearing proposals and instructions affecting field activities; and serving as the central point within the agency through which headquarters offices obtain emergency field support services.
- Providing direction and counsel to Program Directors in the implementation of policies and operational guidelines that form the framework for emergency management of agency field activities.
- Developing and/or recommending to the Commissioner emergency policy, programs, and plans for activities between the agency and State and local governments.
- Administering the agency’s overall Federal-State program and policy and coordinating the program aspects of agency contracts with State and local counterpart agencies.
- Evaluating the overall management and capabilities of the agency’s field organization and initiating action to improve the management of incident-related field activities.
- Maintaining liaison with other Federal agencies and providing assistance to States and localities in the event of a national disaster or other emergencies.
- Participating in emergency preparedness and response activities associated with national events, natural and man-made disasters, and illness outbreaks/injuries.
- Coordinating emergency activities with and providing assistance to Department components and other external stakeholders in the event of a natural disaster or other emergencies.
- Providing planning, monitoring, command and coordination of response to national emergencies and natural and man-made disasters.
- Providing programmatic assistance to States and localities in the event of a natural disaster or other emergency requiring agency assistance.
• Providing oversight of COOP, Safety and local Safety Committee, shelter in place, evacuation plans, and natural disaster management.

**C.5.2.1 Office of Management**

The Office of Management (OM) in an emergency or disaster is responsible for, but not limited to, the following activities:

- Advising the ACRA, other ORA senior managers and staff on all areas of management, including budget formulation and execution, domestic and foreign travel, financial management, human capital management and analysis, ethics, labor relations, safety, and ORA-wide administrative operations and facilities management.
- Overseeing the management of acquisitions and contracts within ORA.
- Adjusting overall manpower allocations based on incident requirements.
- Providing advice, counsel, and support on all areas of labor and employee relations.
- Serving as emergency contact for any issues that occur outside of normal business hours in relation to travel.

**C.5.2.2 Office of Communications and Project Management**

The Office of Communications and Project Management (OCPM) provides, maintains and applies expertise in coordination of executive and legislative correspondence in support of ORA and FDA programs.

OCPM leads the organization’s communications activities and provides counsel and advice to the ORA senior leadership on communications, develops communication strategies and plans to build awareness and promote important organization initiatives, and serves as ORA’s focal point for developing a full range of editorial, web, and digital media information, materials, and products for the organization’s key audiences (consumers, health professionals, industry and employees).

OCPM provides executive secretariat support to ORA including coordinating executive and legislative correspondence and activities including managing and preparing other information such as briefing documents or speeches and presentations.

OCPM is responsible for maintaining FDA’s rapid communication systems that interface with SLTT governments, major municipalities, other Federal partners, and national public health/regulatory associations. Emergency broadcast messages and 50-State conference calls are two examples of ways that information is disseminated to stakeholders. OCPM would work with FDA’s Office of External Affairs to coordinate communication and possibly provide staffing for an FDA JIC.

OCPM may provide personnel in response to an incident to assist in the facilitation of communications with State and local partners. Staff members will be made available around the clock to notify and provide stakeholders with essential and relevant information. If a formal agency response structure such as the IMG is established, OCPM personnel could be assigned to the FDA JIC and perform duties as described in *Section D.2.1.4.6, “State and Local Communications Group”* and *Section E.2.2, “Communicating with States, Terrorities, Tribal Nations, and Local Governments.”* OCPM personnel may also be assigned to the General Staff section, such as Operations, as dictated by the incident response.
C.5.2.3 Office of Criminal Investigations

The Office of Criminal Investigations (OCI) has primary responsibility for all criminal investigations conducted by FDA, including suspected tampering incidents and counterfeit products.\(^{25}\) Additionally, OCI is the primary POC for all law enforcement and intelligence issues pertaining to threats or perceived threats against FDA-regulated products. OCI has special agents at FDA headquarters and in field offices nationwide. OCI also maintains a full-time presence at the National Counterterrorism Center (NCTC), the FBI’s National Joint Terrorism Task Force (NJTTF), and DHS. In case of an emergency or disaster, OCI is responsible for, but not limited to, the following activities:

- Advising and assisting the ACRA and other key FDA officials on regulations, criminal, counter-terrorism, and intelligence matters.
- Maintaining classified electronic connectivity with the U.S. Intelligence Community.
- Directing FDA criminal investigation activities in coordination with other agency components and Federal, State, and local law enforcement agencies.
- Implementing and enforcing agency policy related to criminal investigations.
- Initiating and conducting criminal investigations under all statutes administered by FDA.
- Coordinating assignments and activities involving undercover and surveillance personnel located in field offices throughout the United States.
- Ensuring coordination of criminal investigation activities with FDA District offices and adherence to the agency’s enforcement priorities through cooperative relationships with field and headquarters organizational components.
- Providing recommendations to OCC on referrals of criminal cases to DOJ, or directly to the U.S. Attorney when such direct reference is authorized, for further investigation and/or prosecution.
- Maintaining automated data processing systems to be used for criminal investigations and related enforcement matters.
- Participating in Grand Jury investigations and serving as agents of the Grand Jury.

C.5.2.4 Office of Training Education and Development

The Office of Training Education and Development (OTED) helps design, develop, and deliver the training, as needed. The office works across all ORA program areas. OTED promotes standardized utilization of NIMS according to the FDA NIMS Implementation Plan.

C.5.2.5 Office of Partnerships and Operational Policy

The Office of Partnerships and Operational Policy (OPOP) works collaboratively with State partners on contract inspection programs and implementation of National Standards, such as the Manufactured Food Regulatory Program Standards (MFRPS), as well as cooperative agreements designed to build capability and capacity. OPOP also manages the processes necessary to expedite information sharing through the issuance of commissions that allow the agency to rapidly share non-public information with State and local partners.

OPOP sponsors and provides programmatic/administrative oversight to several funded cooperative agreement programs and grants. This support strengthens State and local food regulatory agency infrastructure

\(^{25}\) Any deliberate contamination of an FDA-regulated product is a criminal act under Title 18 of the U.S.C. Section 1365 (Tampering with Consumer Products).
and enhances our partners’ abilities to effectively respond to emergencies/incidents involving food or feed. Several of these programs, such as the RRT program, involve OP working in collaboration with relevant offices within ORA and OFVM to provide technical guidance and oversight and ensure the programs align with agency policy.

The Office of Strategic Planning and Operational Policy (OSPOP), led by the Assistant Commissioner for Compliance Policy (ACCP), has the primary responsibility for advancing ORA’s goal to be strategic and to fully integrate risk management and data analysis into all policy and resource utilization decisions by:

- Serving as the ORA focal point on significant policy issues that affect ORA programs and providing coordination, development, and direction on such issues.
- Serving as the focal point for testimony requests, information disclosure requests (including those under 21 CFR 20.85, 21 CFR 20.88, and 21 CFR 20.89), and MOUs.
- Conducting risk analysis in support of policy development and work planning and to support the establishment of priorities for inspection, analysis, and compliance actions.
- Establishing and maintaining the ORA work plan in alignment with strategic and risk-based policy decisions.

C.5.2.6 Office of Human and Animal Food Operations

OHAFO coordinates and manages all agency field operations as they relate to the safety of the Nation’s domestically produced and imported human and animal foods, and cosmetics. OHAFO provides oversight of inspectional operations and compliance actions to protect and advance public health. Any part of agency field operations may be called upon to respond to or investigate an emergency. OHAFO develops issues, approves, and clears proposals and instructions affecting field activities; and serves as the central point within the agency through which headquarters offices obtain field support services. The OHAFO offices and their divisions working together with the field handle FDA’s day-to-day and emergency operations throughout the 12 Divisions.

OHAFO coordinates emergency domestic and foreign field investigations. OHAFO represents and makes decisions on behalf of the Assistant Commissioner for Operations (ACO) relative to food/feed program investigatory operations, including emergency response activities and international affairs. OHAFO coordinates and manages all domestic and foreign field investigatory operations related to foods and feed regulated by the agency on behalf of the ACO; serves as the central point within the agency through which directorates and other headquarters offices obtain field support services for food and feed activities; and serves as the agency focal point in coordinating, directing, and assisting the field and headquarters offices with investigative food and feed activities.

OHAFO is composed of Human and Animal Food East, Human and Animal Food West, and State Cooperative Programs, as well as the Audit Staff. The Human and Animal Food West consists of the Division of Domestic Human and Animal Food Operations and the Domestic Produce Safety branches. Human and Animal Food East contains the Division of Foreign Human and Animal Food operations. The Division of Foreign Operations is responsible for, but not limited to, security monitoring of shipments of food for potential contamination with CBRN agents; identifying shipments with potential terrorist connections; and coordinating specific actions against suspect shipments with FDA field personnel, OCI, and DHS CBP.

In an emergency, OHAFO is responsible for, but not limited to, the following activities:

- Providing national or subject matter experts as necessary in an emergency.
- Developing compliance policy and recommending policy to the ACRA.
- Serving as FDA’s focal point for guidance on recall plans and procedures.
• Directing and coordinating field activities in support of all food and animal feed facility investigations and inspections.

• Maintaining liaison with other FDA components, industry, and other government agencies to ensure proper implementation and completion of assessment and investigation of regulated industry, impacted food facilities, and food products.

• Directing multiple Division or Center food and animal feed inspection and investigation activities, which are regulated under the FD&C Act.

C.5.2.7 Office of Medical Products and Tobacco Operations

The Office of Medical Products and Tobacco Operations (OMPTO) represents and makes decisions on behalf of the ACO relative to medical products and tobacco investigatory operations, including emergency response activities and international affairs. OMPTO oversees four offices in the coordination, interpretation, and evaluation of the agency’s overall field inspections and compliance efforts in the areas of medical products and tobacco. OMPTO provides advice and counsel to the ACRA and other senior agency leaders on medical product and tobacco inspection, compliance and other field activities. OMPTO coordinates medical product and tobacco operations with the Office of Enforcement and Import Operations (OEIO) and the Office of Regulatory Science (ORS), and supports medical products and tobacco partnerships and policy through collaboration with the Office of Partnership and Operational and Policy (OPOP). OMPTO also oversees and coordinates across programs medical product and tobacco related recalls, consumer complaints, and quality system activities and directs and coordinates ORA’s emergency preparedness and response activities relative to medical products and tobacco. Each of these offices is led by a Director who directs and oversees the operations of the program in ORA Divisions, as well as ensures coordination between the field and headquarters program functions.

OMPTO Offices:

• Office of Bioresearch Monitoring Operations (OBIMO)

OBIMO includes two divisions, the Division of Bioresearch Monitoring Operations I and II. The Immediate Office of the Director includes the Bioresearch Monitoring Operations Staff. In addition, OBIMO includes Postmarketing Adverse Drug Experience (PADE) and Risk Evaluation and Mitigation Strategies (REMS) inspections.

• Office of Pharmaceutical Quality Operations (OPQO)

OPQO includes the Division of Pharmaceutical Quality Operations I, II, III, and IV; the Division of Pharmaceutical Quality Programs; and the Division of Foreign Pharmaceutical Quality Inspections. Within each of the four field Divisions, there is a Pharmaceutical Quality Investigations Branch and a Pharmaceutical Quality Compliance Branch. (Note: Division 1 includes two Pharmaceutical Quality Investigations Branches.)

• Office of Biological Products Operations (OBPO)

OBPO includes two field divisions, the Division of Biological Products Operations I and II. Within the Immediate Office of the Director are the Biological Products Operations Staff and the Team Biologics Staff.
• **Office of Medical Device and Radiological Health Operations (OMDRHO)**

OMDRHO has three Divisions—the Division of Medical Device and Radiological Health Operations I, II, and III. Within each of the three Divisions, there is a Medical Device and Radiological Health Investigations Branch, and a Medical Device and Radiological Health Compliance Branch. OMDRHO also includes the Foreign Medical Device and Radiological Health Inspection Staff as well as the Medical Device and Radiological Health Operations Staff within the Office of the Directors.

• **Tobacco Operations Staff**

The Tobacco Program includes one cadre of geographically dispersed employees. The Tobacco Operations Staff provides national and foreign coverage to conduct domestic and foreign inspections of tobacco finished product manufacturers, seek compliance and enforcement of the law and regulations designated to reduce the health burden of tobacco use, and conduct investigations at public events where tobacco product manufacturers distribute free samples.

In an emergency, OMPTO is responsible for, but not limited to, the following activities:

- Providing national or subject matter experts as necessary in an emergency.
- Recommending policy to the ACRA.
- Developing, recommending, and reviewing guidance on investigatory procedures related to medical products and tobacco.
- Directing and coordinating emergency preparedness and field response related to medical products and tobacco.
- Coordinating multiple Division inspection and investigation activities and serving as liaison with other FDA components.

**C.5.2.8 Office of Enforcement and Import Operations**

OEIO advises and provides assistance on regulations and compliance matters that impact policy development, implementation, and long-range goals. OEIO coordinates, interprets, and evaluates FDA’s overall compliance efforts within the United States and internationally. This Office acts as liaison with other Federal agencies and international organizations and governments on compliance matters, evaluates proposed legal actions, coordinates these actions with OO and OCC, and handles appeals of proposed compliance actions that are disapproved by the Centers or OCC. OEIO and has responsibilities, which include, but are not limited to, monitoring and controlling import and export activity associated with implicated product(s), countries, foreign manufacturers and shippers, filers, importers, and/or consignees and coordinating surveillance of imported products with the U.S. Customs and Border Protection (CBP).

In an emergency, OEIO is responsible for, but not limited to, the following activities:

- Serving as FDA’s focal point for guidance on recall plans and procedures.
- Directing and coordinating field activities in support of all product recalls and import operations and import-related issues.
- Maintaining liaison with other FDA components, industry, and other government agencies to ensure proper implementation and completion of recall plans and activities.
- Coordinating Center enforcement actions against individuals and companies that violate the FD&C Act.
• Identifying unapproved, fraudulent, harmful, or ineffective FDA-regulated health products and initiating investigations, advisory or enforcement actions, and outreach activities when appropriate.

C.5.2.9 Office of Regulatory Science

ORS reports directly to the ACRA and oversees scientific operations related to human and animal food programs and medical products, and tobacco programs including emergency response activities and international affairs. ORS is ORA’s headquarters scientific office that oversees and manages technical and scientific staff related to human and animal food and medical and tobacco products regulated by the agency.

In an emergency, ORS is responsible for, but not limited to, the following activities:

• Preparing ORA laboratory response and monitoring surveillance databases.
• Activating the Food Emergency Response Network (FERN).
• Developing, recommending, and reviewing guidance on laboratory procedures.
• Coordinating ORA laboratory response.

C.5.2.9.1 Forensic Chemistry Center

The Forensic Chemistry Center (FCC), located in Cincinnati, OH, is part of FDA’s ORA and reports to the Office of Regulatory Science. Its personnel perform forensic analysis of evidence and applied research on a daily basis, regularly providing expert technical support and associated testimony for FDA’s OCI as its primary laboratory. The laboratory also provides forensic analyses of samples in support of all of FDA’s Centers as well as other domestic and foreign public health partners. The FCC plays a major role in many programs initiated by FDA, particularly anti-counterfeiting, anti-tampering, and counterterrorism.

C.5.2.9.2 Winchester Engineering and Analytical Center

The Winchester Engineering and Analytical Center (WEAC) located in Winchester, MA, reports directly to ORA’s Office of Regulatory Science (ORS). WEAC holds ORA’s U.S. Nuclear Regulatory Commission (NRC) Materials License, and its personnel perform radionuclide-related laboratory research and analyses on a day-to-day basis. WEAC serves as FDA’s sole radiation analytical laboratory and provides radioanalytical analyses for samples. WEAC is also ORA’s only device-specialized laboratory and performs testing on various medical devices.

ORA’s Radiation Safety Officer (RSO) is stationed at WEAC. The ORA RSO who reports to ORS’s Office of Business and Safety Operations (OBSO) leads responsibilities for the radiation safety program for ORA laboratories and field offices.
D. DIRECTION, CONTROL, AND COORDINATION

FDA organizes emergency response operations in accordance with the concepts, principles, and terminology of the ICS, as defined within NIMS. Incident Command and subordinate Planning, Operations, Logistics, and Finance/Administration functions lay the foundation for the agency’s implementation of the ICS during an all-hazards response, at headquarters and field levels, and across geographic divides. The inherent design of this system enables rapid, scalable, and flexible agency-wide emergency management activities. Figure D-1 and Figure D-2 depict FDA’s emergency response command and coordinating structures and are followed by a brief description of each designated ICS position.

This section of the FDA EOP describes FDA’s application of the ICS. It is important to remember that two key principles of ICS are its scalability and flexibility. Emergency and disaster situations can be unpredictable and dynamic. To accommodate this, the ICS organizational structure was developed to be expanded easily from a very small size for routine operations to a larger organization capable of handling catastrophic events. Therefore, FDA’s organizational structure, while containing certain core positions, must be tailored to the specific requirements of each incident. Refer to the FDA EOP Incident Annexes for detailed information on FDA organizational constructs for individual hazards.

For individual Center/Office emergency response organizations within this basic structure, refer to each organizational component’s EOP and/or SOPs.

D.1 FIELD INCIDENT COMMAND

The use of Incident Command in the field is achieved through the establishment of an IMT at appropriate field location(s) to determine field incident response objectives and execute the tactical strategy for responding to an incident per the agency’s priorities. The IMT is led by a field IC(s) who operates under the authority of ORA Management via a delegation of authority during an activation and is responsible for the overall management of the incident at the field level. ORA Management may be represented by a District Director (DD) or a Program Executive Group (PEG) consisting of the DD and the relevant Program Division Directors (PDDs) of the programs impacted by the incident. Depending on the scope and scale of the incident, the IMT will coordinate with a headquarters (HQ) coordinating structure such as an Incident Management Group (IMG) (see Section B.2.2.3.3, “Headquarters Response”), the Coordinated Outbreak and Response Evaluation Network (CORE), or OEO. An IMG and IMT may or may not mobilize and/or demobilize concurrently.

An IMT functions in accordance with ICS principles; it is scalable and flexible based on the complexity of the incident. The IC may establish Operations, Planning, Logistics, and/or Finance/Administration Sections with subordinate positions, as needed, to manage the agency’s on-the-ground response. Figure D-1 is an example of how an IMT may be organized.

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26 For more information on the ICS, refer to www.training.fema.gov/EMIWeb/IS/ICSResource/index.htm.
27 Depending on the level and complexity of incident requirements, certain agency ICS functions and positions may not be necessary. Refer to the FDA EOP Incident Annexes for specific personnel assignments.
The IC will direct the field personnel to conduct tactical operations to determine the cause/source of the emergency and/or resolve the emergency, based on the IAP they develop and, in some cases, information, analysis, and direction received from the IMG. Depending on the nature and stage of the emergency, these operations might include facility or field inspections, sample collection, laboratory analysis, traceback and traceforward investigations, import-related activities, recall efforts, and enforcement actions. In conducting these operations, Incident Command might interact with regulated entities, FDA headquarters, State and local officials, other Federal agencies, foreign regulatory authorities, and regional/local news media.28

When appropriate, the FDA field IC may be part of a UC with multiple ICs representing different agencies (e.g., FDA and a State health agency) or jurisdictions (e.g., Federal, State, local). The field IC may also be requested to provide a Liaison Officer (LNO) to a Regional or local UC.

28 The IMG Public Information Officer coordinates development and release of agency public health and consumer protection messages for use with the public, media, and/or other agencies with approval from FDA headquarters and HHS.
D.1.1 Command Staff

The command staff consists of the Liaison Officer, Safety Officer and Public Information Officer. When assigned, they report directly to the Incident Commander and provide information, safety, and liaison services in support of the incident.29

D.1.1.1 Incident Commander

The IC is responsible for the overall management of the incident and for providing direction and guidance to the Command and General Staffs. The IC assigns the Operations, Planning, Logistics and Finance/Administration Section chiefs as needed. When these positions are not filled, the IC is responsible for accomplishing or managing these aspects of the incident organization. The IC must assess the overall requirements of the incident and determine the best course of action for the incident management team to pursue. The key responsibilities of the IC are to establish priorities, determine incident objectives and direction for managing the incident, and to approve and authorize the implementation of the IAP.

D.1.1.2 Liaison Officer

For incidents that may be multijurisdictional, a LNO may be assigned as the point of contact for supporting agencies. The LNO’s primary function is to establish and coordinate interagency contacts and to keep the agencies informed of the incident’s status.

D.1.1.3 Safety Officer

The Safety Officer monitors incident operations and advises the IC on all matters relating to operational safety. The SO has the authority to immediately alter, delay, suspend, or terminate any and all operations that pose a danger to the life and health of personnel.

D.1.1.4 Public Information Officer

FDA has specific policies for handling public information. The IMT PIO works with the appropriate FDA organizational components to coordinate message development and dissemination as applicable. The IMT PIO maintains an accurate account of information about the incident to inform and update management, stakeholders, and the IMG PIO or JIC (if activated).

D.1.2 General Staff29

The General Staff is responsible for the major functional aspects of the IMT. The General Staff consists of Operations, Planning, Logistics, and Finance/Administration Sections. Each Section is led by a Section Chief who reports directly to the IC.

D.1.2.1 Operations Section

The Operations Section is responsible for managing field level tactical operations to meet the objectives set forth by the IC. Operations can be organized and executed by Strike Teams, Task Forces, Single Resources, Divisions, Groups, and/or Branches depending on the size, scope, and complexity of the incident.

29 For a full description of IMT Command and General Staff responsibilities, refer to FDA’s Incident Management Handbook (IMH).
D.1.2.2 Planning Section

The Planning Section is responsible for collecting, evaluating, and disseminating information pertaining to the incident. This Section documents and maintains information on the current and forecasted situation, as well as the status of FDA resources assigned to the incident. The Planning Section prepares the IAP, develops SitReps, organizes meetings and teleconferences, and maintains the status of resources assigned to the incident. The Section may include up to four primary units: Resources, Situation, Demobilization, and Documentation.

D.1.2.3 Logistics Section

The Logistics Section is responsible for all service and support requirements of the team. This Section provides facilities, transportation, security, supplies, communications, and IT support services to field personnel as required.

D.1.2.4 Finance/Administration Section

The Finance/Administration Section is responsible for all financial considerations associated with an incident. This Section verifies time, approves procurements, manages compensation/claims, and tracks the overall cost of the response.

D.2 Headquarters Incident Coordination

When necessary, headquarters will activate an incident coordination structure referred to as the IMG. (See Section B.2.2.3.3, “FDA Headquarters Response,” regarding which FDA officials have the authority to activate an IMG.) The IMG facilitates the integration of all FDA assets responding to an incident as required by the NRF. As the agency’s coordination structure, the IMG serves as the focal point for headquarters-level strategic coordination and provides recommendations for actions to take and priorities for the use of resources in support of incident management. The IMG has general oversight of the application of FDA resources in coordination with existing agency and interagency resource management. It provides strategic situational awareness and decision support across the full spectrum of incident management domains—prevention, protection, mitigation, response, and recovery.

Because of FDA’s unique organizational structure, which involves emergency response resources and capabilities from multiple Centers and Offices, the FDA IMG coordinates unified agency responses to incidents, which is somewhat similar to an Area Command. The goal of the IMG is to facilitate a comprehensive, integrated, and coordinated approach to domestic and international incident management.

For its coordination structure, FDA may adapt the headquarters-based Incident Command structure to address the scope of the incident, which at times acts as a hybrid coordination and command structure, as in the case of a pandemic influenza response. Figure D-2 is an example of how an IMG may be organized. This structure is flexible, and parts of it may be expanded or contracted as the emergency evolves.
The FDA OEM may request that the FDA product Centers and OC Offices provide staff to support emergency operations based on the subject matter expertise required. FDA product Centers and OC Office staff may hold any position in the IMG (Command or General Staff) as requested by the AIC.

**D.2.1  Command Staff**

**D.2.1.1  Agency Incident Coordinator**

The AIC has overall delegated authority and responsibility for agency-wide emergency operations and ensures information review for appropriate action, including referral to the Commissioner or AEG for executive-level direction or to headquarters or field units for any necessary follow-up. In addition, the AIC serves as the central point for headquarters coordination regarding issues and decisions involving FDA organizational components, the HHS SOC, other HHS OPDIVs and Staff Divisions, applicable Federal and State agencies, and foreign governments. The AIC is responsible for assigning the Operations, Planning, and other Section chiefs or, when these additional positions are not required, personally accomplishing or managing these aspects of the incident organization. If an IMG is activated, the AIC and designated support staff will be physically located in the EOC, unless weather, building emergencies, or some other unforeseen circumstance prevents it.

Until the Commissioner; Deputy Commissioner; or other designated senior agency official such as the Director, OSEM, appoints an AIC for any emergency or disaster involving multiple FDA organizational components and/or requiring external coordination, the OEM/OEO Director will serve as the AIC and will report to the Commissioner or his/her authorized designee. He/she is responsible for coordinating and providing situational awareness on all incident activities performed at the headquarters and field levels.

The AIC responsibilities are to:

- Develop broad objectives for agency-wide emergency response operations.
- Coordinate with engaged headquarters and field Centers/Offices in the development of individual incident objectives and strategies.

30 Refer to the *FDA Incident Management Handbook* for the specific time-phased emergency actions of the AIC.
- Allocate/reallocate agency resources as the established priorities change.
- Ensure effective communications between headquarters and field organizational components.
- Ensure overarching incident management objectives are met and do not conflict with each other or with agency policies.
- Identify and report on critical agency resource needs.
- Facilitate FDA’s transition to full recovery operations.
- Establish communications channels with any field Incident Command structure (e.g., IMT).
- Ensure that a formal incident After Action Review is completed and recorded after the incident stabilizes.

The FDA Command Staff of the IMG and possibly in the field includes a Public Information Officer (PIO) and LNOs who report directly to the AIC. Additional Command Staff positions may be required, depending on the nature, scope, complexity, and number of locations of the incident(s), or according to specific requirements established by the AIC. At headquarters, a Legal Advisor may be assigned to the Command Staff as a Technical Specialist or to the Planning Section to advise the AIC on legal matters, such as emergency proclamations, and the legality of EUAs (legal rights and restrictions pertaining to product use). Similarly, a Medical Advisor or Science Advisor may be designated and assigned directly to the Command Staff to provide advice and recommendations to the AIC in the context of incidents involving medical and public health response or medical countermeasures considerations. Command Staff positions are discussed in the paragraphs that follow.31

**D.2.1.2 Liaison Officers**

LNOs interface or report from external Federal agencies (i.e., HHS, CDC, USDA, EPA, DHS, FBI, or DOS) as representation to FDA during an incident. These Command Staff members serve as direct links between the AIC and their agency, providing input on their agencies’ policies, activities, concerns, resource availability, and other incident-related matters. These LNOs are responsible for maintaining an awareness of the situation, reporting relevant information, and facilitating communication regarding FDA requests for assistance from the LNO’s agency. Staff assigned to these positions must have the authority to speak for their parent agencies or organizations on all matters, following appropriate consultations with their agency leadership. Assistants and personnel from other agencies or organizations, public or private, involved in incident management activities may be assigned to the LNO to facilitate coordination. There may also be a need for a liaison with industry.

**D.2.1.3 Safety Officer**

The Safety Officer monitors incident operations and advises the AIC on all matters relating to operational safety, including the health and safety of emergency responder personnel. The Safety Officer is, in turn, responsible to the AIC for the systems and procedures necessary to ensure ongoing assessment of hazardous environments, including the incident safety plan, coordination of multiagency safety efforts, and implementation of measures to promote emergency responder safety as well as the general safety of incident operations. The Safety Officer has immediate authority to stop and/or prevent unsafe acts during incident operations. It is important to note that the agencies, organizations, or jurisdictions that contribute to joint safety management efforts do not lose their individual identities or responsibility for their own programs, policies, and personnel. Rather, each contributes to the overall effort to protect all responder personnel involved in incident operations.

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31 Refer to the *FDA Incident Management Handbook* for specific time-phased emergency actions of the Command Staff.
D.2.1.4 Public Information Officer

The FDA JIC (see Section D.2.1.4.1) will be led by the IMG PIO and Deputy IMG PIO. The IMG PIO works with the appropriate FDA organizational components such as the product Centers and OEA to coordinate message development and dissemination. FDA has specific policies for handling public information. The IMG PIO major responsibilities generally apply to any type of incident involving an FDA-regulated product. Additional staff may be tasked to support the IMG PIO.

The FDA IMG PIO maintains regular communications with OEA as well as other Center and Office (i.e., OIP) public affairs and communications staffs and informs the IMG of any associated public information activities, concerns, and information requests. The IMG PIO is not necessarily the incident spokesperson, but the coordinator of message development and dissemination.

In addition to the headquarters role, there may be one or more field IMT PIOs. The IMT PIO maintains an accurate account of information about the incident to inform and update the IMG PIO. For FDA incidents, the on-scene PIO must coordinate all information and media interactions with the IMG PIO.

The relationship between these entities is illustrated in Figure D-3. Proper media clearance channels through HHS are not superseded by the figure below. Dependent on the incident’s strategic communication objectives, the IMG PIO’s responsibilities may be shared among other individuals and Offices in a FDA JIC as part of a larger Joint Information System (JIS).

Figure D-3. FDA’s Joint Information System (JIS)

The FDA JIC staffing composition and structure are determined by the various methods FDA uses to reach its multiple audiences. The following structure (Figure D-4) depicts the groups and liaisons that could be part of a FDA JIC.

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32 For more information, refer to the *FDA Joint Information Center Handbook*.

33 **Joint Information System**: Integrates incident information and public affairs into a cohesive organization designed to provide consistent, coordinated, accurate, accessible, timely, and complete information during crisis or incident operations. The mission of the JIS is to provide a structure and system for developing and delivering coordinated interagency messages; developing, recommending, and executing public information plans and strategies on behalf of the IC; advising the IC concerning public affairs issues that could affect a response effort; and controlling rumors and inaccurate information that could undermine public confidence in the emergency response effort.
Figure D-4. Possible Structure of FDA Joint Information Center

The majority of incidents to which FDA responds do not call for all eight reporting elements depicted in the FDA JIC figure above. The four groups are the elements most likely to be activated. When span of control becomes an issue for the IMG PIO, the IMG Deputy PIO may be assigned to manage the FDA JIC staff.

D.2.1.4.1 FDA Joint Information Center

FDA’s JIC is a physical collocated group of representatives from the FDA Offices and Centers involved in the event who are designated to handle public information needs. The FDA JIC monitors and assesses the need for and effectiveness of public messaging regarding the incident and FDA’s response. The FDA IMG PIO works with the FDA JIC to provide a public messaging strategy for the incident. The AIC and AEG (see Section B.2.2.3.3 “FDA Headquarters Response”) approve the strategy and messages. Target audiences include consumers, industry, Congress, and health professionals.

The FDA JIC will be established in proximity to FDA’s EOC to provide a base of operation for the public information function to facilitate timely communication among its members. If a physical location within FDA facilities is not possible, the FDA JIC may function as a group using electronic communications such as conference calls, emails, and instant messaging. The FDA JIC will be activated when the IMG PIO and the AIC deem it will benefit the incident objectives to have the communication strategies coordinated in the same physical location or by electronic communications.

An interagency unified JIC may be established at the national level to ensure consistent and accurate public messages are issued when multiple agencies and organizations are involved in a response. An interagency JIC is a collocated or collaborating group of representatives designated by their agencies and organizations to participate in the joint development, coordination, and delivery of public messages. An interagency JIC ensures the coordinated release of information under two NRF annexes—ESF #15 – External Affairs Annex and the Public Affairs Support Annex. If FDA is asked to participate in an interagency JIC, the IMG PIO will identify someone to serve as FDA’s representative on the interagency JIC. This representative will serve as a liaison to FDA’s JIC and report to the IMG PIO.
D.2.1.4.2 Media Relations Group
During an incident response, the Associate Commissioner for External Affairs, in conjunction with the Director of OEA’s OMA, will appoint an experienced staff media officer to lead the FDA JIC’s Media Relations Group with responsibility for managing the news media outreach activities and message development of the FDA JIC.

D.2.1.4.3 Web and Digital Media Group
During an incident response, the Associate Commissioner for External Affairs, in consultation with the Director of OEA’s Web and Digital Media staff, will appoint a lead for the FDA JIC Web Group to manage the agency’s web and digital communications related to the incident, including web postings (e.g., portals, response questions and answers [Q&As]), FDA Voice blog, Twitter, Facebook, Flickr, widgets, searchable databases (for extensive recalls), postings on www.foodsafety.gov, or other government-wide websites such as www.recalls.gov.

D.2.1.4.4 Stakeholder Outreach Group
During an incident response, the Associate Commissioner for External Affairs, in conjunction with the Director of OEA’s Office of Health and Constituent Affairs (OHCA), will appoint a lead for the Stakeholder Outreach Group to manage outreach to and message development for external stakeholders, as part of the FDA JIC. As needed, representatives from other OC Offices or Centers will be added to the group. For example, OEA’s Office of Communications will frequently be involved in advising on consumer messages and preparing external communications materials such as Consumer Updates, FDA Voice blogs, and FDA Fact Sheets.

D.2.1.4.5 State and Local Officials Group
During an incident response, the Director of ORA’s OP, in consultation with the Associate Commissioner for External Affairs, will appoint a lead for the State and Local Officials Group to manage the outreach and two-way communications with State and local officials as part of the FDA JIC.

D.2.1.4.6 Call Centers Liaison
During an incident response, the Associate Commissioner for External Affairs, in consultation with Directors of OEM and Center/Office communications offices, will appoint a Call Centers Liaison to the FDA JIC to ensure that up-to-date messages related to the incident and response are available to the general public and industry through the relevant agency Call Centers (if needed).

D.2.1.4.7 Congressional Liaison
During an incident response, the Associate Commissioner for Legislation, in conjunction with the Associate Commissioner for External Affairs, will appoint a Congressional Liaison to manage communications with congressional members as part of the responsibilities of the FDA JIC (if needed).

D.2.1.4.8 International Liaison to External Unified JIC
If an incident includes an international dimension, the Associate Commissioner for International Programs, in conjunction with the Associate Commissioner for External Affairs, will appoint an International Liaison who will manage communications with international partners as part of the FDA JIC (if needed).

34 For FDA, the term stakeholder includes industry and industry associations, consumers and patients, health professionals, scientists and researchers, and the general public, among others.
D.2.1.4.9 Internal Communications Liaison

During an incident response, the Associate Commissioner for External Affairs, in consultation with OEA’s Assistant Commissioner for Communications and the Associate Commissioner for Operations, will appoint an Internal Communications Liaison to ensure that FDA employees receive up-to-date information about the incident and the agency’s response efforts (if needed).

D.2.2 General Staff

The General Staff is responsible for the functional aspects of any ICS. The General Staff consists of Operations, Planning, Logistics, and Finance/Administration Sections composed of designated FDA headquarters and field emergency staff. The responsibilities associated with these Sections are discussed more fully below.35

D.2.2.1 Operations Section

The Operations Section is responsible for coordinating all agency activities in response to an identified threat or hazard, establishing an agency-wide common operating picture, and restoring normal operations. This Section may be composed of a Section Chief, Divisions, and Groups (each led by a designated Supervisor). Figure D-5 depicts FDA’s Operations Section.

![Figure D-5. Operations Section Organizational Elements](image)

Expansion or contraction of the Operations Section may vary according to numerous considerations and operational factors associated with an incident. In some cases, a functional approach may be used. In other cases, the organizational structure will be determined by geographical or Center/Office jurisdictional boundaries or a mix of functional and geographical considerations may be appropriate. The AIC will determine the appropriate organizational structure, based on previous experience and the specific circumstances of the incident at hand.

D.2.2.1.1 Operations Section Chief

Until the AIC appoints an Operations Section Chief, an emergency coordinator from OEM/OEO designated by the AIC will fulfill the role. The Operations Section Chief is responsible to the AIC for managing incident activities required to implement the agency IAP, as well as providing the AIC with regular status reports. The Operations Section Chief monitors initial and ongoing headquarters and field response activities, advises and assists the AIC on appropriate courses of action, responds to requests for assistance from internal organizational components and external partners, and coordinates regular status/situation updates throughout the lifecycle of the incident. He/she has direct involvement in providing input to and implementing the IAP and serves as the focal point for intra- and interagency coordination and communications. The Operations Section Chief is collocated with the AIC at the FDA EOC.

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35 Refer to the FDA Incident Management Handbook for specific time-phased emergency actions of the General Staff.
D.2.2.1.2 Divisions and Groups

Divisions may be established within the Operations Section to assign responsibility based on the physical/geographic location(s) of an emergency or major disaster. Groups may also be assembled to combine similar activities being addressed for a single or multiple incident(s). For example, geographic Divisions may be created in support of a single or multiple responding FDA Districts, or to coordinate operations when a foreign country is involved. Functional Groups, on the other hand, may include personnel addressing investigations, enforcement, traceback, laboratory analysis, and other issues, depending on the nature and scope of the incident. Divisions and Groups are staffed by representatives from the OC, ORA, and Centers and may change both in designation and composition based on the specific incident requirements. They typically operate from the FDA EOC or alternate workspace locations as needed. A Supervisor shall be designated for each to oversee operations and provide status/situation reporting to the Operations Section Chief.36

D.2.2.1.3 Center/Office Emergency Coordinators

FDA product Centers and OC Offices shall designate a senior staff member and alternates to serve as the Center/Office Emergency Coordinator. The Center/Office Emergency Coordinator is responsible for advising the Center/Office Director of actions needed to address an incident, supervising and reporting on the Center’s/Office’s emergency response activities, triaging information requests received from the IMG Command or General Staff to appropriate Center/Office SMEs, and channeling necessary communications between the Center and other FDA organizational components (e.g., other Centers/Offices and the FDA EOC).

He/she serves as the focal point for internal and external Center/Office communications during emergencies and disasters and is responsible for providing regular status reports as requested. During some large-scale or catastrophic incidents, Center/Office emergency coordinators may be directed to report to the FDA EOC to support initial or prolonged emergency operations.

D.2.2.2 Planning Section

The Planning Section is responsible for collecting, evaluating, and disseminating information pertaining to the incident. This Section documents and maintains information on the current and forecasted situation, as well as the status of FDA resources assigned to the incident. The Planning Section prepares the IMG IAP, compiles and consolidates FDA-wide SitReps, and organizes status/situation reporting meetings and teleconferences. As shown in Figure D-6, the Planning Section, led by a Section Chief, may include subordinate units (i.e., Resources, Situation, Documentation, and/or Demobilization) and Technical Specialists from FDA headquarters Centers and Offices to assist in evaluating the situation and forecasting requirements for additional agency resource support.

36 Refer to the Incident Annexes of this EOP for detailed information on the functional groups established within the Operations Section to address specific incident requirements.
Figure D-6. Planning Section Organizational Elements

D.2.2.2.1 Planning Section Chief

Until the AIC appoints a Planning Section Chief, an emergency coordinator from OEM/OEO designated by the AIC will fulfill the role. This individual oversees all data gathering and analysis activities regarding incident operations and assigned resources, coordinates planning and status/situation reporting meetings, and assists in the preparation and consolidation of the agency-level IAP. The Planning Section Chief is collocated with the AIC and Operations Section Chief at the FDA EOC. The Planning Section Chief is responsible for supervising the preparation and distribution of SitReps. See Section E.1.2, “Situation Reporting,” for additional information on SitReps.

D.2.2.2.2 Units

The Planning Section may have up to five primary units (described below) that support the Section Chief. Generally composed of headquarters staff, each unit operates from the FDA EOC unless otherwise directed.

- **Resource Unit.** Makes certain that FDA resources (i.e., personnel, teams, and equipment) are available for assignment to or deployment during an incident. During IMG activations, the Planning Section Chief will generally activate a Status/Check-In Recorder under the Resource Unit, who will maintain an accurate record of all personnel reporting to and demobilizing from the IMG.

- **Situation Unit.** Collects, processes, analyzes, and organizes ongoing situation information; prepares situation summaries; and develops projections and forecasts of future events related to the incident.

- **Documentation Unit.** Maintains a record of the major steps taken to resolve the incident and files incident records for legal, analytical, and historical purposes.

- **Demobilization Unit.** Develops an Incident Demobilization Plan that includes specific instructions for agency resources to return to normal operations.

- **Intelligence Unit.** The intelligence function can be used to provide information that leads to the determination of cause, projection of spread, assessment of impact, or selection of countermeasures for a given incident (regardless of the source) such as public health events and disease outbreaks of unknown origins. In the later context, intelligence includes not only national

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37 The incident itself will primarily dictate the need for Planning Section units. Refer to the Incident Annexes of this EOP for detailed information on the types and composition of units needed for each situation.
security or other types of classified information, but also other operational information such as risk assessments and medical intelligence (i.e., surveillance). It must be analyzed and shared with appropriate personnel, designated by the Agency Incident Coordinator (AIC), to assist in decision making.

D.2.2.2.3 Technical Specialists

Technical Specialists have specialized skills and are activated only when needed. Technical Specialists are most often assigned to the specific area (e.g., Section, Branch, Unit, Division) where their services are needed and performed. These headquarters personnel may be assigned to the Planning Section, but may support any ICS function, including the Command Staff working directly for the AIC. Technical Specialists normally perform the same duties during an incident that they perform in their everyday jobs and operate from normal locations unless directed to report to the FDA EOC by the Planning Section Chief or AIC.

The following are examples of the kinds of specialized expertise that may be required during an emergency or disaster:

- **Geographic Information Specialist.** Collects, analyses, and disseminates location-based information to support the objectives of the IMG. Common roles include identifying areas affected by the incident and locating FDA assets and regulated industry. The Geographic Information Specialist produces maps and data products for use in situational awareness, operations management, and reporting. Requests for maps should be submitted to the Geographic Information Specialist. The Geographic Information Specialist position is normally located under the Planning Section but may assist other Sections of the headquarters IMG and field IMT as necessary.

- **Legal Specialist.** Usually is a Manager or other Senior Attorney in the OCC or in OCET who provides advice on issues that involve application or interpretation of relevant statutes and regulations. For example, in some situations involving possibly adulterated food, the Legal Specialist might advise the IMG on the legal aspects of requesting records from food facilities under Section 414 of the FD&C Act. As another example, the Legal Specialist also reviews any EUA requests that might be submitted to the agency regarding medical countermeasures needed to protect responders and the public. In addition, the Legal Specialist considers legal issues in media releases and other documents prepared for the incident response.

- **Science/Medical Specialist.** Advises on scientific and/or medical issues that arise during the response. The Science/Medical Specialist will have expertise in the scientific and/or medical aspects of the incident and will typically be a staff member from the lead Center(s) for the emergency.

- **International Specialist.** Typically a staff member from OIP who advises on international issues that arise during the response.

- **Product Interface Specialist.** A key Center SME, depending on the medical products necessary for the response, and an OCET staff member may advise on medical countermeasure issues that arise during the response. An OCET staff member may lead the coordination of cross-agency activities related to medical countermeasure use and availability activities.

As needed, other Technical Specialists with expert knowledge in the following areas may be required:

- Human/veterinary medicine
- Pharmacology
- Statistics

38 The incident itself will primarily dictate the need for Technical Specialists. Refer to the FDA EOP Incident Annexes for detailed information on the types and numbers of Technical Specialists needed for each situation.
D.2.2.3 Logistics Section

The Logistics Section is responsible for all service support requirements needed to facilitate effective and efficient agency incident management operations. This Section provides facilities, transportation, security, supplies, communications, and IT support services to FDA headquarters and field personnel as required. The Logistics Section is led by a Section Chief. This role is initially filled by an OEM Emergency Coordinator or staff member; unless designated otherwise by the AIC, support units may be staffed from the Office of Management (OM), OIMT, and other headquarters Offices as appropriate.

Figure D-8. Logistics Section Organizational Elements

D.2.2.3.1 Units

Units, such as Supply, Facilities, Communications, and Services Acquisition (described below), may be established within the Logistics Section based on the scope and complexity of incident operations. Composed of headquarters staff, each unit operates from normal work locations unless otherwise directed. The Logistics Section Chief will determine, in coordination with the AIC and given current and anticipated requirements, the need for establishing specific subordinate units.

- **Supply Unit.** Coordinates ordering, receiving, processing, storage, inventory management, and distribution of supplies and materials for headquarters personnel.
- **Facilities Unit.** Coordinates headquarters facility operations and building security support services during incident operations.
- **Communications Unit.** Coordinates headquarters IT systems and equipment in support of incident operations and supervises and operates the Communications Center.
- **Services Acquisition Unit.** Coordinates requirements for contracted services needed during the emergency response.

D.2.2.3.2 Technical Specialists

Technical Specialists have specialized skills and are activated only when needed. These headquarters personnel are assigned to the Logistic Section, but may support any ICS function. Technical Specialists normally perform the same duties during an incident that they perform in their everyday jobs and operate from normal locations unless directed to report to the FDA EOC by the Logistics Section Chief or AIC. The OEM EOC Manager is an example of a Technical Specialist who may be required during an emergency to provide specialized expertise.

D.2.2.3.3 EOC Manager

The EOC Manager will assist the IMG in performing the emergency response operations, which requires that EOC communication capabilities be functioning at all times, including access to network, databases, and the Internet. The EOC Manager is responsible for the technical requirements during COOP relocations and the operation of the EOC in accordance with the OEM COOP Plan.
In addition, the EOC Manager may assist with the transmission of electronic information during an emergency and analysis of information/data needs for a response. Onsite, the EOC Manager coordinates all enhancements of the agency’s data information systems, which pertain to critical and time-sensitive information.

**D.2.2.4 Finance/Administration Section**

A Finance/Administration Section is established only for those incidents requiring large-scale or extended agency operations or specific financial and administrative support services. Some of the functions that fall within the scope of this Section are accounting, payment processing, financial reporting, foreign and domestic travel expenditures, employee relocation, payroll liaison, and financial systems management. The Finance/Administration Section is led by a Section Chief staffed with support personnel as appropriate.

![Figure D-9. Finance/Administration Section Organizational Elements](image)

**D.2.2.4.1 Units**

Units, such as Compensation and Claims, Procurement, Cost, and Time (described below), may be established within the Finance/Administration Section based on the scope and complexity of incident operations. Composed of headquarters staff, each unit operates from normal work locations unless otherwise directed. The Finance/Administration Section Chief will determine, given current and anticipated requirements, the need for establishing specific subordinate units.

- **Compensation and Claims Unit.** Handles agency injury/illness compensation and claims.
- **Procurement Unit.** Administers all financial matters pertaining to vendor contracts and is modified as needed.
- **Cost Unit.** Provides cost analysis data for the incident.
- **Time Unit.** Ensures proper daily recording of personnel time in accordance with FDA policies.

**D.2.3 Agency Executives**

**D.2.3.1 Agency Executive Group**

The AEG is established when an emergency requires the involvement of senior FDA officials with the knowledge and authority to address a wide range of policy and resource issues. The AEG serves primarily to provide strategic policy direction and guidance for major agency emergency response activities and to approve important policy decisions, in consultation with the Commissioner and AIC. Decisions that may be within the AEG’s scope include addressing unprecedented legal, regulatory, policy, and resource

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39 In certain incidents, the Logistics and Finance/Administration Sections may be collapsed together under one Branch. During an influenza pandemic or alert, it is likely FDA’s ICS organizational structure will expand to include Logistics and Finance/Administrative as part of the response.
problems and impasses brought about by the emergency and that impact the agency’s ability to respond effectively.

Typically, members of the AEG include:

- OSEM Director or designee
- OEM Director or Designee
- ORA Associate Commissioner or Deputy Associate Commissioner
- ORA Office Director, for Program-specific incidents
- OEA Associate Commissioner
- Senior leadership from appropriate Offices/Centers (i.e., OCS/OCET, OCC, OGROP/OIP, OFVM, OPP, and OO/OFBA/OB) or their representatives

**D.2.3.2 FDA Executive Committee**

The FDA Executive Committee is a standing organizational management structure implemented by the Commissioner that addresses cross-cutting issues requiring the input of key agency leaders. The Committee employs the strength of a cross-cutting leadership structure to facilitate the governance of FDA. Issues in the following areas are identified and addressed:

- Operational issues
- Strategic direction and priorities
- Scientific and policy issues

The Commissioner serves as Chairperson, unless delegated to the Principal Deputy Commissioner, a Deputy Commissioner, the Chief of Staff, or the Chief Operating Officer. Committee members are:

- Principal Deputy Commissioner
- Deputy Commissioners
- Chief Operating Officer
- Chief Counsel
- Chief of Staff
- Directors of CBER, CDER, CDRH, CFSAN, CTP, CVM
- Associate Commissioner for Regulatory Affairs
- Chief Scientist

Other participants and observers who are full-time or permanent part-time employees of the agency or other such employees of the Federal government may participate and attend in certain circumstances. Such invitees should be confirmed by the Chairperson and/or designated staff and should generally attend the meeting only in relation to an agenda item.

**D.2.4 Multilagency Coordination**

Many of the incidents to which FDA responds involve other agencies that have a stake in providing response or needed resources. If the product involved in an incident crosses agency jurisdictional lines,
the Multiagency Coordination (MAC) system may be used to support a unified coordination of operations. The primary function of the MAC system is to coordinate activities above the field level and to prioritize incident demands for critical or competing resources, thereby assisting the coordination of field operations.

In this unified coordination approach, representatives from each agency meet to set goals and decide how each agency can contribute to the achievement of the goals. There can be strong, formal command and control relationships between and among the agencies, or the command and control linkages can be based on informal but structured arrangements that recognize Federal and State responsibilities. It can be as simple as a teleconference or, alternately, require an assembled group and associated support systems. Such an assembled group comprises senior officials from the agencies involved in the response who are brought together to form a MAC Group, providing executive guidance to individual agency Incident Command groups on policy, resource allocation, and communications.

A MAC Group may be supported by a MAC Group Coordinator, who may supervise MAC Group Situation Assessment and Resources Information Units that collect and assemble information needed for the MAC Group to fulfill its mission. These units would obtain such information from the agency’s IMG/IMT. The MAC Group may also have its own Public Information Unit to coordinate summary information and access to information sources with the media and other governmental entities. This function is often called a Joint Information Center (or JIC).

The results of the MAC Group’s deliberations are distributed by its members directly to their own organizations as well as through the normal chain of command (e.g., EOCs, Incident Command/Coordination Groups, etc.).

Regardless of the level of government involved in response to an incident, all MAC Groups have five key functions. These functions include:

1. **Direction and Control.** Provide indirect control and direction for complex or multijurisdictional incidents. Serve as a single POC for prioritizing incidents and for facilitating access to critical resources.
2. **Information Collection and Evaluation.** Serve as a central point for collecting, analyzing, and interpreting information from a variety of sources.
3. **Coordination.** Play a key role in coordinating the information flow and resources for complex incidents or multiple incidents occurring simultaneously.
4. **Priority Setting.** Prioritize incidents and critical resources using the priorities established by the NPG as well as the priorities used to guide development of incident objectives (Life Safety, Incident Stabilization, and Property and Environmental Conservation), and use these priorities at the policy level.
5. **Resource Management.** Manage scarce resources in line with incident priorities; resource management includes identifying and acquiring needed resources in addition to allocating existing or known resources.

**D.2.5 Multiagency Coordination Group for Foodborne Illness Outbreaks**

FDA, along with USDA agencies, HHS, and the CDC, has created a Multiagency Coordination Group for Foodborne Illness Outbreaks (MAC-FIO) (see Figure D-9). This approach to unified coordination includes a MAC Group that comprises senior officials from each agency, and these personnel provide

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40 Information excerpted from FACT SHEET: Multiagency Coordination Group for Foodborne Illness Outbreaks (MAC-FIO), November 2012.
executive guidance to individual agency Incident Command groups on policy, resource allocation, and communications.

**Figure D-10. MAC-FIO Unified Coordination for Foodborne Illness Outbreaks**

The MAC-FIO may be alerted when indicators of a large-scale foodborne illness outbreak potentially involving more than one agency become evident. This pre-alert is intended to ensure the critical coordination among agencies involved under a unified structure for all large-scale foodborne illness outbreaks involving multiple agencies. A formal MAC-FIO in place can also be activated expeditiously to provide the executive guidance to Incident Command through the agencies’ response coordinating bodies and ensure:

- Coordination of the development and implementation of a flexible, scalable, and adaptable response among Federal, State, local, and tribal governments to multijurisdictional foodborne illness outbreaks.
- Coordination between law enforcement agencies and public health agencies of investigations of foodborne illness outbreaks due to intentional contamination.
- Coordinated communication strategy to the public, industry, and other international and domestic partners.
- Coordinated collaborative efforts among Federal, State, and local governments in identifying the source of the contamination and removal of contaminated products/ingredients through a coordinated traceback and recall of all contaminated food, feed, and ingredients.
- Coordinated development of measures to prevent future contamination and illnesses.
- Established systematic and proactive approach for managing incidents across multiple jurisdictions as well as established a national-level policy coordination structure.
E. COMMUNICATIONS AND INFORMATION MANAGEMENT

Effective emergency management and incident response activities rely upon flexible communications and information systems that provide for a “common operating picture” across all agency components and staff. Properly planned, established, and used communications processes and systems enable the vertical and horizontal dissemination of information among and between FDA organizational components; with Federal, SLTT, and foreign government agencies; and to the news media, industry, and consumers.

E.1 INTRA-AGENCY COORDINATION

During the course of any incident for which an FDA response may be necessary, timely communications and information sharing among agency units is critical to assist FDA leadership and staff with gaining and maintaining situational awareness and making decisions. Incident information, such as emergency alerts and status/situation reporting, can aid in developing an agency IAP, serve as the basis for releasing public messages, determine the need for involvement of FDA organizational components, and satisfy RFIs. The following are examples of information generated by responsible FDA emergency staff and the technological systems they rely upon for use during an incident to build an agency-wide “common operating picture.”

E.1.1 Emergency Alerts

FDA may be alerted to a threat or hazard through a variety of means, including from FDA headquarters Centers/Offices or District offices, other government agencies, consumers, and industry. Formal notifications typically occur by phone (primary) to the OEM/OEO with appropriate follow-up by email and/or fax referencing the initial emergency alert.

An OEM/OEO Emergency Coordinator records the initial call and subsequent follow-up information. After regular duty hours (8:00 a.m. to 4:30 p.m. Eastern Time, Monday through Friday, excluding Federal holidays) and when the Office is not operating, an answering service refers all emergency calls to an OEO Late Duty Officer (LDO). Upon receipt, the Emergency Coordinator or LDO uses a predeveloped call-down tree/notification list for internal and external distribution of emergency alerts.

For each incident, OEM/OEO will provide notification about the incident to those appropriate officials and staff of the Centers and Offices who may have a response role in that particular incident.
Also, the OEM/OEO Director may send notification of EOC operating status level and additional staffing requirements to all involved parties.

### E.1.2 Situation Reporting

Standardized incident reporting and documentation procedures ensure that agency-wide situational awareness is maintained and provide emergency personnel with easy access to critical information. Status reports relay information specifically related to the availability or assignment of agency resources. SitReps offer a snapshot of each tasked organizational component’s emergency operations and contain confirmed information regarding the explicit details of the incident (who, what, when, where, and how). Transmission of this data in a common format, and at predesignated intervals, enables FDA to rapidly share critical intelligence and information between headquarters and field elements and with external partners as appropriate.

During an incident, FDA IMTs, field offices, and Centers may provide regular SitReps via the EON-IMS, which is discussed in-depth below. These SitReps highlight significant event information and emergency response activities (e.g., investigations, analyses, public affairs, cooperating agencies, scientific data, legal court matters). District, Senior, and Center emergency coordinators are responsible for configuring the SitRep process, defining recipients, and reviewing and accepting inputted information.

Once field SitReps have been approved/confirmed by emergency coordinators, the Planning Section Chief, in coordination with Command and General Staff located at the FDA EOC, assembles a consolidated SitRep and posts to the EON-IMS. This report combines relevant data from field SitReps for dissemination across FDA and is used to provide operational information to external Federal agencies.

### E.1.3 Emergency Operations Network – Incident Management System

In addition to primary modes of voice and data communications (telephone and email), the Emergency Operations Network - Incident Management System (EON-IMS) serves as the central hub for exchanging, storing, and relaying all incident-related information within the agency. Managed by OEM, this system integrates multiple data streams from other electronic systems, such as the Electronic Submission Gateway (ESG), the EON Call Center, the Mission Accomplishment and Regulatory Compliance Service-Recall Enterprise System (MARCS-RES), and from FDA laboratories and investigators and external agencies, into a coherent fashion during critical decision points. The EON-IMS creates a safety net that significantly reduces the probability that incidents will prevent FDA from accomplishing its objectives and minimizes the impact of these events on normal operations. It provides a web-based connection for all agency organizational components and their partners through which accurate real-time data about an incident can be shared and discussed, and is used in all situations requiring efficient receipt and dissemination of large volumes of information to FDA stakeholders, including the public and other government agencies.

The EON-IMS, which is critical for comprehensive performance of agency incident management functions, has three components: (1) incident tracking and contact management, (2) a collaboration and knowledge management tool for meetings and document management, and (3) a GIS for mapping and impact assessment. Through EON-IMS, FDA ensures that its emergency response is uniform, consistent, and coordinated. Participants are able to provide input and access real-time data, agency emergency plans and procedures, contact databases, and analysis tools, which enhance response capabilities.

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41 Additional EON-IMS resources (e.g., user manual, reference sheets, etc.) may be found on the FDA intranet at http://inside.fda.gov:9003/ProgramsInitiatives/EmergencyPreparednessandResponse/EmergencyOperationsNetworkIncidentManagementSystemEONIMSResources/default.htm or by contacting OEM.
E.1.4 Government Emergency Telecommunications Service

In response to White House guidance, the National Communications System (NCS) developed the Government Emergency Telecommunications Service (GETS) to ensure key personnel can communicate over local and long-distance telephone networks during an emergency. GETS can provide users a high likelihood of call completion during the most severe conditions of network congestion and disruption. There is an additional feature that can be activated on the GETS card called Wireless Priority Service (WPS). WPS allows GETS users to have their official cell phone number associated with their GETS service and ensures cellular communications are also available. Located on each GETS card are instructions for its use.

E.2 INTERGOVERNMENTAL COORDINATION

Coordination with responsible government agencies at the Federal and SLTT levels must be effective during emergency and disaster situations to ensure that resource allocations are efficient, policy is understood, and that roles are well defined. Considering that Federal agency responsibility varies from one type of incident to another, and that State and local government organizations differ from those of the Federal government, the specific number of agencies that should cooperate in a given situation is highly dependent on the incident at-hand and its location(s) as well. However, specific FDA organizational components are charged with establishing and maintaining communications at the various intergovernmental levels, regardless of incident size, complexity, or geographic location. These headquarters and field organizations are described below.

E.2.1 Communicating with Federal Government Agencies

OEM/OEO staff is responsible for coordinating Federal interagency liaison activities and establishing communications with the headquarters emergency operations offices and staff of Federal agencies during incidents of varying nature and scope. Through the use of dedicated communications channels, standardized reporting mechanisms, and the deployment of internal and external liaisons, OEM/OEO is able to maintain contact with the HHS SOC, CDC Director’s Emergency Operations Center (DEOC), USDA/FSIS, and other Federal department/agency EOCs as appropriate. OEM/OEO or the IMG represents the agency on national level conference calls, with input from IMTs, Centers, and Offices as needed. Centers may have technical consultation with other government agencies during an emergency and provide updates about significant consults to the agency through the IMG if activated, or OEM/OEO.

The lead District, in coordination with the FDA EOC, is also responsible for communicating with appropriate Federal field offices and incident management field structures, such as the DHS/FEMA Regional Response Coordination Center (RRCC), JFO, and any established disaster recovery centers.

During emergency response operations, a requirement may exist to conduct secure communications with other government agencies. This situation requires strict adherence to guidelines and regulations for handling, processing, and storing sensitive/classified materials. FDA restricts access to sensitive and classified material to ensure its release to only those agency personnel who possess appropriate security clearances and a need-to-know. OSEM/OSO maintains a list of locations and numbers of agency secure voice and data equipment. This list is available at:


E.2.2 Communicating with State, Local, Tribal, and Territorial Governments

During an incident, both the lead District and other investigating Districts are responsible for establishing communications with responsible State agencies. Usually, FDA will work through the State in coordinating efforts at the local level or through the use of the State’s predesignated RRTs. However, depending upon the State, it may be more appropriate for FDA District offices, in coordination with the FDA EOC, to work directly with local government to address the situation.
The Office of Communications and Quality Program Management (OCQPM) maintains a rapid communication system to alert officials in State governments, major municipalities, and poison control centers and to quickly enlist nationwide assistance for the agency’s emergency operations. During an incident, the OCQPM will:

- Ensure the appropriate State agencies, such as agriculture and health departments, are notified of significant confirmed incidents within their States or which cross State borders, indicating the potential for problem FDA-regulated products to enter commerce.
- Prepare (or distribute) information requested by States during the emergency response operations and ensure they are fully advised as to what action(s) FDA can recommend under the circumstances of the specific incident.

### E.2.3 Communicating with Foreign Governments

During an incident when commerce with Canada or Mexico is involved, FDA’s response coordination will be performed by OEM/OEO in cooperation with OIP. When commerce with other countries is involved, OIP, in direct coordination with OEM/OEO, will establish, supervise, and/or coordinate and maintain appropriate communication channels.

### E.3 Public Messaging

Public messaging consists of the agency processes, procedures, and systems to communicate timely, accurate, and accessible public health and medical information to the media and directly to FDA stakeholders (both directly and indirectly affected). Information must be coordinated and integrated across FDA organizational components at the headquarters and field levels, as well as with other Federal and SLTT government agencies and industry partners to protect consumer well-being and decrease the risk of illness, injury, disease, or death. Well-developed communications plans and strategies executed by responsible FDA staff help to ensure that messages, alerts and warnings, educational materials, and situational updates are developed and distributed to numerous audiences in a timely, consistent manner.

FDA’s OMA, within the OEA, is responsible for coordinating the release of information to the public during an emergency or disaster. It serves as the agency’s primary liaison with the news media and develops FDA’s public health and consumer protection messages. During an emergency, OMA Public Affairs Specialists, in direct coordination with the assigned IMG PIO, Center media representatives, and all responding field offices, issue press releases, media advisories, and other public statements; respond to media requests; and arrange and support media interviews.

OEA’s Office of Communications develops and promotes FDA consumer health information, which consists of time-sensitive public health and product safety alerts and information about FDA’s roles and responsibilities in protecting and promoting public health. The goals of this Office during an incident include promoting improved outreach and readership of FDA consumer health information, sharing time-sensitive public health and product safety alerts, and facilitating public understanding of FDA’s roles and responsibilities in protecting and promoting public health.

OEA’s Web and Digital Media staff ensures that vital and important information and consumer health messages are expeditiously posted to FDA’s website (www.fda.gov) and updated as needed. The Web and Digital Media staff also coordinates web communications with OMA, HHS/ Assistant Secretary for Public Affairs (ASPA), other Federal agencies, and State and local governments to ensure a consistent and comprehensive approach that reduces duplication of messages.
E.3.1 Coordination with Other Federal Agencies

Coordination with other external Federal agencies is also conducted to ensure the provision of a common message. In accordance with the NRF, pre-identified incident communications protocols are established and ready for use during an incident requiring a coordinated Federal response.

The National Incident Communications Conference Line (NICCL) is used for transmission and exchange of critical and timely (e.g., “breaking”) incident information among Federal and affected SLTT authorities. If the nature of the incident is of critical importance and urgency, DHS Public Affairs will maintain a controller on the line continuously to provide and receive updates from departments and agencies. During sustained incident management activity, the NICCL will be used for daily or other incident communications coordination calls. DHS Public Affairs will maintain a summary of key NICCL communications and interagency coordination actions.

Throughout an incident, OMA will take guidance from and work closely with the HHS Office of the ASPA in publicizing public health messages and implementing a strategic communications plan that:

- Coordinates and expedites FDA messages with HHS/ASPA and other partners to ensure consistency.
- Disseminates clear and accurate messages that maintain, increase, or restore public trust and confidence.
- Uses a variety of vehicles to distribute public health messages.

OEA also coordinates web communications with HHS/ASPA, other Federal agencies, and State and local governments to ensure a consistent and comprehensive approach that reduces duplication of messages.

E.3.2 Congress

FDA’s OL directs and manages FDA’s legislative needs, pending legislation, oversight activities, and congressional relations consistent with the mission of the agency. In an emergency or disaster, the OL is responsible for, but not limited to, the following activities:

- Keeping Congress apprised of FDA actions relative to the incident in coordination with the HHS Office of the Assistant Secretary for Legislation (ASL).
- Responding to congressional RFIs related to the incident.
- Arranging and supporting congressional meetings, briefings, and hearings.
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F. AUTHORITIES AND REFERENCES

The legal authorities that guide the structure, development, and implementation of the FDA EOP include statutes and regulations, presidential directives, national strategies, and Federal government plans and guidance. In addition, internal emergency plans and procedures augment the FDA EOP and provide specific guidance to agency personnel during emergencies and disasters. These documents are listed chronologically by category and summarized below.42

Authorities and references specific to individual hazards are also included in the FDA EOP Incident Annexes.

F.1 STATUTES AND REGULATIONS

F.1.1 Federal Food, Drug, and Cosmetic Act


a. In consultation with experts in science, medicine, and public health and, in cooperation with consumers, users, manufacturers, importers, packers, distributors, and retailers of regulated products, FDA is charged with (1) promoting the public health by promptly and efficiently reviewing clinical research and taking appropriate action on the marketing of regulated products in a timely manner, and (2) with respect to such products, protecting the public health by ensuring that foods are safe, wholesome, sanitary, and properly labeled; human and veterinary drugs are safe and effective; there is reasonable assurance of the safety and effectiveness of devices intended for human use; cosmetics are safe and properly labeled; the public health and safety are protected from electronic product radiation; and tobacco products are properly labeled and are not contaminated.

b. FDA is charged with ensuring that safe and effective drugs, biologics, and devices are available for emergency use.

c. If FDA finds a reasonable possibility that a tobacco product contains a manufacturing or other defect not ordinarily contained in tobacco products on the market that would cause serious, adverse health consequences or death, the agency can order that distribution of any such tobacco product immediately cease and can order a recall as necessary. FDA may also provide notice to persons subject to the risks associated with the use of such tobacco product.

d. If FDA determines that a tobacco product presents an unreasonable risk of substantial harm to the public health, the agency can provide notice to the public as necessary to eliminate such risk, including issuing public service announcements.

e. FDA is authorized to inspect the records of each person who manufactures, processes, transports, distributes, receives, holds, packages, exports, or imports tobacco products if the agency has a

42 This list is not exhaustive, and the associated summaries should not be used as a substitute for the authorities and references themselves.
reasonable belief that a tobacco product is part of illicit trade, smuggling, or counterfeiting operations.

f. The Commissioner of Food and Drugs or designee may request a firm to initiate a recall when a product has been distributed that presents a risk of illness or injury or gross consumer deception; when the responsible firm has not initiated a recall of the product; and when an agency action is necessary to protect the public health and welfare. A request by FDA that a firm recall a product is reserved for urgent situations and is to be directed to the firm that has primary responsibility for the manufacture and marketing of the product that is to be recalled.

g. Under the Project BioShield Act of 2004 (P.L. 108-276), the FD&C Act is amended to allow FDA, in response to a declaration issued by the HHS Secretary, to authorize the use, with appropriate safeguards, of unapproved drugs, devices, or biological products or the use of approved products for unapproved uses. Before FDA can issue an EUA, the HHS Secretary must declare an emergency justifying the authorization to use the product, based on one of three determinations: (1) a determination by the Secretary of Homeland Security of a domestic emergency, or the significant potential for a domestic emergency; (2) a determination by the Secretary of Defense of a military emergency, or the significant potential for a military emergency; (3) a determination by the HHS Secretary of a public health emergency under PHSA or , (4) the Secretary of Homeland Security makes a Material Threat Determination.

h. The Pandemic and All-Hazards Preparedness Act (P.L. 109-417), December 2006, amends the FD&C Act to direct FDA to establish a team of experts on manufacturing and regulatory activities (including compliance with current GMP) to provide both offsite and onsite technical assistance to the manufacturers of qualified countermeasures, security countermeasures, or vaccines at the request of such a manufacturer and at the discretion of the Secretary, if the Secretary determines that a shortage or potential shortage may occur in the United States in the supply of such vaccines or countermeasures and that the provision of such assistance would be beneficial in helping alleviate or avert such shortage.

i. The Pandemic and All-Hazards Preparedness Reauthorization Act (PAHPRA), (P.L. 113-5), March 2013, builds on work the U.S. Department of Health and Human Services has undertaken to advance national health security, amending the Public Health Service Act, authorizing funding through 2018 for buying medical countermeasures under the Project BioShield Act, and increasing the flexibility of BioShield to support advanced research and development of potential medical countermeasures. PAHPRA also enhances the authority of the U.S. Food and Drug Administration to support rapid responses to public health emergencies. The purpose of the Pandemic and All-Hazards Preparedness Reauthorization Act is “to reauthorize certain programs under the Public Health Service Act and the Federal Food, Drug, and Cosmetic Act with respect to public health security and all-hazards preparedness.”

The Food Safety Modernization Act of 2011 (FSMA) (P.L. 111-353) amends the FD&C Act as follows:

a. If FDA reasonably believes 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 must permit FDA to have access to and copy certain records relating to such article and to any other article of food the Secretary reasonably believes is likely to be affected in a similar manner.
b. If FDA believes 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 must permit FDA to have access to and copy certain records relating to such article and to any other article of food the Secretary reasonably believes is likely to be affected in a similar manner.

c. FDA may order the detention of any article of food that is found during an inspection, examination, or investigation if FDA has reason to believe that such article is adulterated or misbranded.

d. If FDA determines there is a reasonable probability that an article of food (other than infant formula) is adulterated under Section 402 of the FD&C Act or misbranded under Section 403 of the FD&C Act and the use of or exposure to such article will cause serious adverse health consequences or death to humans or animals, the Secretary must provide the responsible party with an opportunity to cease distribution and recall such article. If the responsible party refuses to or does not voluntarily cease distribution or recall such article within the time and in the manner prescribed by the Secretary, the Secretary may, by order, require such person to (A) immediately cease distribution of such article; and (B) as applicable, immediately notify all persons (i) manufacturing, processing, packing, transporting, distributing, receiving, holding, or importing and selling such article; and (ii) to which such article has been distributed, transported, or sold, to immediately cease distribution of such article.

e. Food facilities must report to FDA as soon as practicable, but in no case later than 24 hours, within determining there is a reasonable probability that the use of, or exposure to, an article of food manufactured, processed, packed, or held at the facility will cause serious adverse health consequences or death to humans or animals.

f. If FDA determines that food manufactured, processed, packed, received, or held by a facility has a reasonable probability of causing serious adverse health consequences or death to humans or animals, FDA may suspend the registration of a facility that was responsible for such reasonable probability, or knew of or had reason to know of such reasonable probability, and packed, received, or held such food. If a facility’s registration is suspended, no person can introduce food from such facility into interstate or intrastate commerce in the United States.

F.1.2 Public Health Service Act

The Public Health Service Act (PHSA), (42 U.S.C. § 262): National Preparedness for Bioterrorism and Other Public Health Emergencies, as amended, charges FDA with assisting the Secretary, other HHS OPDIVs, and Federal and SLTT government partners in response to bioterrorism and other public health emergencies.43

F.1.3 Federal Anti-Tampering Act

The Federal Anti-Tampering Act (FATA), 18 U.S.C. 1365, authorizes FDA to investigate any tampering of FDA-regulated consumer products.43

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43 To find these Public Laws, go to https://congress.gov/ and click on Public Laws.
F.1.4 Bioterrorism Act

FDA is responsible for carrying out certain provisions of the Public Health Security and Bioterrorism Preparedness and Response Act of 2002 (Bioterrorism Act) (P.L. 107-188).43 Particularly, Subtitle A (Protection of Food Supply) and Subtitle B (Protection of Drug Supply) of Title III provide for the following FDA authorizations:

- FDA may detain any article of food found during an inspection, examination, or investigation that presents a threat of serious adverse health consequences or death to humans or animals.
- FDA is authorized to access certain records when there is a reasonable belief that an article of food is adulterated and presents a threat of adverse health consequences or death to humans or animals.
- Owners and operators of foreign or domestic food facilities that manufacture or process, pack, or hold food for human or animal consumption in the United States must submit information (i.e., identity of the food, manufacturer and shipper, grower, country of origin, country from which the food is shipped, and anticipated port of arrival) to FDA about the facility and emergency contacts.
- FDA shall receive prior notice of imported food shipments before the food arrives at any U.S. port, which must include the article, the manufacturer and shipper, the grower (if known within the specified time in which notice is required), the country of origin, the country from which the article is shipped, and the anticipated port of arrival, and conduct a bioterrorism risk assessment.
- FDA is charged with providing for research on tests and sampling methodologies designed to test food to detect adulteration rapidly, particularly methodologies that detect intentional adulteration and tests that are suitable for inspections of food at ports of entry to the United States.
- FDA is authorized to conduct examinations and investigations through the officers and employees of another Federal department or agency, pursuant to an MOU, at facilities or other locations that are jointly regulated by FDA and such department or agency. FDA is required to notify States when there is credible evidence or information indicating that a shipment, or portions of a shipment, of imported food presents a threat of serious adverse health consequences or death to humans or animals.
- Another Federal department’s or agency’s officers and employees are authorized to conduct examinations and investigations on FDA’s behalf, pursuant to the signing of an MOU between FDA and the head of the other Federal agency.
- FDA, in direct coordination with the CDC and USDA, is charged with coordinating the surveillance of zoonotic diseases.

F.1.5 Stafford Act

F.1.6 Public Readiness and Emergency Preparedness Act

The Public Readiness and Emergency Preparedness Act of 2005 (PREP Act) (P.L. 109-148), PHSA § 319F-3 and § 319F-4, authorizes the Secretary to issue a declaration to provide immunity from tort liability for certain claims that are causally related to development, distribution, administration, or use of “covered countermeasures” and authorizes an emergency fund in the U.S. Department of the Treasury for compensation for injuries from covered countermeasures (www.phe.gov/Preparedness/legal/prepact/Pages/default.aspx).

F.1.7 Title 18 and Title 21 United States Code

FDA investigative jurisdiction includes the FD&C Act and other related Acts (Title 21 U.S.C. and related Acts), FATA (Title 18 U.S.C. § 1365) and other criminal violations of Title 18 U.S.C.4343

F.1.8 Code of Federal Regulations Title 21

Title 21, Chapter I, is the portion of the CFR that governs FDA-regulated products within the United States. These regulations are enforced by FDA, based on the FD&C Act and other applicable laws (www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfctfr/cfbegin.cfm).
PRESIDENTIAL DIRECTIVES

The following presidential directives mandate action from all Federal departments and agencies, including FDA as an HHS OPDIV.

F.1.9 PPD-2, National Strategy for Countering Biological Threats

This Strategy outlines the vision for addressing the challenges from proliferation of biological weapons or their use by terrorists (https://www.hsdl.org/?abstract&did=31404).

F.1.10 PPD-8, National Preparedness

Presidential Policy Directive 8 (PPD-8), National Preparedness, requires the development of a national preparedness goal and national preparedness system. PPD-8 is designed to facilitate an integrated, all-of-Nation/whole community, capabilities-based approach to preparedness.


F.1.11 National Preparedness Goal

The National Preparedness Goal (NPG) sets the vision for nationwide preparedness and identifies the core capabilities and targets necessary to achieve preparedness across five mission areas laid out under PPD-8: Prevention, Protection, Mitigation, Response, and Recovery (www.fema.gov/learn-about-presidential-policy-directive-8).

F.1.12 National Preparedness System

The National Preparedness System is an integrated set of guidance, programs, and processes, enabling the Nation to meet the NPG. Designed to guide domestic efforts of all levels of government, the private and nonprofit sectors, and the public, the National Preparedness System includes guidance for planning, organization, equipment, training, and exercises needed to build and maintain domestic capabilities in support of the NPG (www.fema.gov/learn-about-presidential-policy-directive-8).

F.1.13 National Planning Frameworks

The National Planning Frameworks, which are part of the National Preparedness System, set the strategy and doctrine for building, sustaining, and delivering the core capabilities identified in the NPG. They describe the coordinating structures and alignment of key roles and responsibilities for the whole community and are integrated to ensure interoperability across all mission areas (www.fema.gov/learn-about-presidential-policy-directive-8).

There is one framework for each of the five preparedness mission areas addressed in PPD-8:

- National Prevention Framework
- National Protection Framework
- National Mitigation Framework
- National Response Framework (second edition)
- National Disaster Recovery Framework
F.1.14 **HSPD-5, Management of Domestic Incidents**

HSPD-5 requires the Secretary of Homeland Security to develop and administer a national incident management system that will provide a consistent nationwide approach for Federal, State, and local governments to work effectively and efficiently together to prepare for, respond to, and recover from domestic incidents, regardless of cause, size, or complexity. HSPD-5 also requires the Secretary of Homeland Security, after consultation with appropriate Federal officials, to develop and administer a National Response Plan (NRP). The Directive orders the heads of all Federal agencies to adopt NIMS and NRP and to provide assistance to the Secretary in their development and maintenance (www.dhs.gov/laws-regulations-1).

F.1.15 **HSPD-7, Critical Infrastructure Identification, Prioritization, and Protection**

HSPD-7 establishes a national policy for Federal agencies to identify and prioritize U.S. critical infrastructure and key resources and to protect them from terrorist attacks (www.dhs.gov/xabout/laws/ge_1214597989952.shtm).

F.1.16 **HSPD-9, Defense of United States Agriculture and Food**

HSPD-9 establishes a national policy to defend the agriculture and food system against terrorist attacks, major disasters, and other emergencies (https://www.hsdl.org/?view&did=444013).

F.1.17 **HSPD-18, Medical Countermeasures Against Weapons of Mass Destruction**

HSPD-18 establishes an approach for the development and acquisition of medical countermeasures for attacks involving CBRN agents (https://www.hsdl.org/?view&did=456436).

F.1.18 **NSPD-51/HSPD-20, National Continuity Policy**

This directive, also known as National Security Presidential Directive 51 (NSPD-51), establishes a comprehensive national policy on the continuity of Federal government structures and operations and a single National Continuity Coordinator responsible for coordinating the development and implementation of Federal continuity policies (https://www.hsdl.org/?view&did=476323).

F.1.19 **HSPD-21, Public Health and Medical Preparedness**

HSPD-21 establishes a national strategy that will enable a level of public health and medical preparedness sufficient to address a range of possible disasters (https://www.hsdl.org/?view&did=480002).

F.1.20 **HSPD-22, Domestic Chemical Defense**

HSPD-22 establishes a national policy and directs actions to strengthen the ability of the United States to prevent, protect, respond to, and recover from terrorist attacks employing toxic chemicals and other chemical incidents. (This directive is classified and, thus, not publicly available.)
F.2 NATIONAL STRATEGIES

F.2.1 National Strategy for the Physical Protection of Critical Infrastructures and Key Assets

The National Strategy for the Physical Protection of Critical Infrastructures and Key Assets establishes a foundation for building and fostering the cooperative environment in which government, industry, and private citizens can carry out their respective protection responsibilities more effectively and efficiently. Moreover, this Strategy identifies a clear set of national goals and objectives and outlines the guiding principles that underpin the efforts to secure the infrastructures and assets vital to national security, governance, public health and safety, economy, and public confidence (www.dhs.gov/national-strategy-physical-protection-critical-infrastructure-and-key-assetsNational Strategy for Physical Protection of Critical Infrastructure and Key Assets | Homeland Security).

F.2.2 National Strategy for Homeland Security

The National Strategy for Homeland Security guides, organizes, and unifies the Nation’s homeland security efforts. Homeland security is a responsibility shared across the entire Nation, and the Strategy provides a common framework to prevent and disrupt terrorist attacks; protect the American people, critical infrastructure, and key resources; respond to and recover from incidents that do occur; and continue to strengthen the foundation to ensure long-term success. It builds upon the 2002 version; reflects an increased understanding of the terrorist threats confronting the United States today; incorporates lessons learned from exercises and real-world catastrophes, including Hurricane Katrina; and proposes new initiatives and approaches that will enable the Nation to achieve its homeland security objectives (www.dhs.gov/xabout/history/gc_1193938363680.shtm).

F.2.3 National Security Strategy


F.3 FEDERAL PLANS AND GUIDANCE

F.3.1 Comprehensive Preparedness Guide 101

The Comprehensive Preparedness Guide (CPG) 101: Developing and Maintaining Emergency Operations Plans expands on FEMA’s efforts to provide guidance about response and recovery planning to SLTT governments. It also extends those planning concepts into the prevention and protection mission areas. CPG 101 integrates concepts from the National Preparedness Guidelines, NIMS, NRF, National Strategy for Information Sharing (NSIS), and National Infrastructure Protection Plan (NIPP), and it incorporates recommendations from the 2005 Nationwide Plan Review (https://www.fema.gov/media-library/assets/documents/25975).

F.3.2 National Incident Management System

NIMS provides a systematic, proactive approach to guide departments and agencies at all levels of government, NGOs, and the private sector to work seamlessly to prevent, protect against, respond to, recover from, and mitigate the effects of incidents, regardless of cause, size, location, or complexity, to
reduce the loss of life and property and harm to the environment. NIMS works hand-in-hand with the NRF. It provides the template for the management of incidents, while the NRF provides the structure and mechanisms for national-level policy for incident management.

NIMS is not an operational incident management or resource allocation plan. It represents a core set of doctrines, concepts, principles, terminology, and organizational processes that enable effective, efficient, and collaborative incident management (https://www.fema.gov/nims-doctrine-supporting-guides-tools).

NIMS has established ICS as the standardized incident organizational structure for the management of all incidents.

**F.3.3 National Infrastructure Protection Plan**

NIPP, and its supporting Sector-Specific Plans (SSPs), provides a coordinated approach to critical infrastructure and key resources (CIKR) protection roles and responsibilities for Federal and SLTT government and private sector security partners. NIPP sets national priorities, goals, and requirements for effective distribution of funding and resources, which help ensure the Nation’s government, economy, and public services continue in the event of a terrorist attack or other disaster (www.dhs.gov/xlibrary/assets/NIPP_Plan_noApps.pdf).

The SSPs provide the means by which the NIPP is implemented across all 18 CIKR sectors, as well as a national framework for each sector to address its unique characteristics and risk landscape. FDA is responsible for the food sector (except meat; poultry; and frozen, dried, and liquid eggs, which are under the authority of the USDA/FSIS). Agency food sector protection efforts are detailed in the Agriculture and Food CIKR SSP to the NIPP (www.fda.gov/downloads/Food/FoodDefense/ucm081308.pdf). FDA also supports HHS, which is responsible for the SSP for the public health and medical sector.

**F.3.4 National Preparedness Guidelines**

The National Preparedness Guidelines organizes and synchronizes national (including Federal and SLTT) efforts to strengthen preparedness. The Guidelines guide national investments in national preparedness, incorporate lessons learned from past emergencies and disasters into national preparedness priorities, facilitate a capability- and risk-based investment planning process, and establish readiness metrics to measure progress and a system for assessing the Nation’s overall preparedness capability to respond to major events, especially those involving acts of terrorism. There are four critical elements to the Guidelines: (1) the National Preparedness Vision; (2) the 15 National Planning Scenarios; (3) the Universal Task List (UTL); and (4) the Target Capabilities List (TCL) (www.dhs.gov/national-preparedness-guidelines).

**F.3.5 National Response Framework**

The NRF presents the guiding principles that enable first responders, decision makers, and supporting entities to prepare for and provide a unified national response to emergencies and major disasters—from the smallest incident to the largest catastrophe. It establishes a comprehensive, national, all-hazards approach to domestic incident management and defines the key principles, roles, and structures that organize the way the Nation responds.

The NRF, an update to the 2004 NRP, describes how communities, tribes, territories, States, the Federal government, and private sector and nongovernmental partners apply these principles for a coordinated, effective national response. This document also identifies special circumstances where the Federal government exercises a larger role, including incidents where Federal interests are involved and catastrophic incidents where a State would require significant support. The NRF enables first responders, decision makers, and supporting entities to provide a unified national response (https://www.fema.gov/media-library/assets/documents/117791).
F.3.6 National Disaster Recovery Framework

The NDRF (https://www.fema.gov/national-disaster-recovery-framework) is a guide that enables effective recovery support to disaster-impacted SLTT jurisdictions. It provides a flexible structure that enables disaster recovery managers to operate in a unified and collaborative manner. It also focuses on how best to restore, redevelop, and revitalize the health, social, economic, natural, and environmental fabric of the community and build a more resilient Nation.

The NDRF is consistent with the vision set forth in PPD-8 (https://www.fema.gov/learn-about-presidential-policy-directive-8), National Preparedness, which directs FEMA to work with interagency partners to publish a recovery framework.

The NDRF introduces six new Recovery Support Functions (https://www.fema.gov/recovery-support-functions) that provide a structure to facilitate problem-solving, improve access to resources, and foster coordination among State and Federal agencies, nongovernmental partners, and stakeholders. Each RSF has coordinating and primary Federal agencies and supporting organizations that operate together with State, local, and tribal government officials, NGOs, and private sector partners.

The NDRF presents three positions that provide focal points for incorporating recovery considerations into the decision-making process and monitoring the need for adjustments in assistance where necessary and feasible throughout the recovery process. Those positions are Federal Disaster Recovery Coordinator (FDRC), State or Tribal Disaster Recovery Coordinators (SDRCs or TDRCs), and Local Disaster Recovery Managers (LDRMs) (www.fema.gov/national-disaster-recovery-framework).

F.4 FDA SUPPORTING DOCUMENTATION

F.4.1 Center/Office Emergency Plans and Procedures

Some FDA organizational components have developed individual emergency plans and operating procedures that define the scope of incident management activities necessary for that Center or Office. These EOPs and subordinate procedural documents present specific details on Center/Office response to emergencies and disasters, such as assigning staff roles, responsibilities, and lines of authority; detailing projected times, places, and coordination mechanisms for carrying out emergency actions; describing how staffs and facilities are protected; and identifying the personnel, equipment, facilities, supplies, and other resources necessary for use during response and recovery operations.

The following represent the compendium of existing FDA emergency/disaster plans and procedures. Each supports implementation of the FDA EOP and is available to assist FDA personnel during prevention, preparedness, protection, mitigation, response, and recovery activities.

- FDA Incident Management Handbook (September 2013). This handbook is designed to assist FDA personnel in the use of ICS and NIMS during incident response operations and planned events. The handbook is an easy reference job aid for responders and includes position-specific checklists. The handbook provides guidance and protocols for responders and management to understand what their positional responsibilities are under ICS/NIMS and how these positions integrate with the overall response structure.
- FDA Joint Information Center Handbook (November 2013). This handbook is for personnel assigned to a FDA JIC as part of an IMG incident response or planned event. The handbook
describes the organization of FDA personnel tasked with public information responsibilities. The handbook provides an organized approach to providing accurate, coordinated, timely, and accessible information to FDA audiences as an emergency situation evolves.


Copies of individual Center and Office emergency plans and procedures may be found on the Center/Office intranet web pages on *inside.fda*.

**F.4.2 SOP for FDA Headquarters Building Closures and Dismissal of Employees**

The current operational status of FDA headquarters’ buildings and dismissal of employees related to administrative situations including all FDA locations in the Washington, D.C., metropolitan area, including the White Oak Campus, is provided at [www.fda.gov/AboutFDA/WorkingatFDA/BuildingsandFacilities/default.htm](http://www.fda.gov/AboutFDA/WorkingatFDA/BuildingsandFacilities/default.htm).

**F.4.3 Center/Office COOP Plans**

FDA has designed COOP Plans for each of its headquarters and field organizational components, including OC, ORA, CBER, CDER, CDRH, CFSAN, CVM, NCTR and District offices. These plans ensure that agency essential functions can continue to be performed from alternate locations during all hazards, including localized acts of nature, accidents, and technological or attack-related situations.
G. PLAN MANAGEMENT

G.1 COORDINATION

The FDA OEM is responsible for the overall management and maintenance of the FDA EOP (including all annexes and appendices) and for ensuring that changes and revisions occur, including their preparation, coordination, publication, and distribution. OEM, in consultation with all responsible agency Centers and Offices, coordinates plan reviews to address lessons learned from real-world and simulated incidents and incorporates necessary organizational or technological changes.

G.2 MAINTENANCE

Any FDA headquarters or field organizational component may propose changes to the FDA EOP. All proposed changes are to be captured within the “Record of Changes” section to this EOP and submitted to OEM for formal review. Approval from the appropriate Center/Office senior official should be secured prior to submitting any proposed change.

Once OEM receives a proposed change to the FDA EOP, it will coordinate a review with all applicable Centers and Offices to determine its suitability for inclusion in the plan. Once this coordination has been finalized, OEM will issue an official Notice of Change agency-wide and provide the intra- and Internet links to the updated plan.

- The Notice of Change will specify the date, number, subject, purpose, background, and action required, and provide the change language.
- Once published, the changes will be considered part of the FDA EOP for operational purposes pending a formal revision and re-issuance of the entire document.

G.3 PROMULGATION

The OEM Director is responsible for coordinating reviews of the basic plan and all applicable annexes to the FDA EOP and for re-adoption and distribution every four years or more frequently as deemed necessary. The annual review will consider lessons learned and best practices identified during exercises and/or actual emergencies and disasters. Any changes that result from reviews will be incorporated in the FDA EOP and any supporting Center/Office emergency plans and procedures.

Prior to any reissuance of the EOP, or at the request of the Commissioner or his/her authorized representative, OEM will work with agency headquarters and field emergency coordinators and others as appropriate to revise key areas of the plan. OEM will disseminate the final draft of the revised FDA EOP for review and concurrence prior to submitting to the Commissioner for approval and promulgation.

G.4 STANDARDS FOR SUPPORTING DOCUMENTATION

The FDA EOP, including the basic plan and all annexes, is the core plan for FDA emergency operations and provides the structures and processes for coordinating agency incident management activities for emergencies, including natural disasters, terrorist attacks, and other criminal acts. It provides an umbrella configuration for existing emergency plans and procedural documents (with appropriate modifications and revisions) as integrated components, supplements, or supporting material.

All FDA organizational components must incorporate common principles, concepts, and language when developing or updating individual emergency plans and procedures. Accordingly, all Center/Office emergency documentation shall be compatible with the FDA EOP.
G.5 TRAINING AND EXERCISES

FDA shall conduct and/or participate in HHS and other organizations’ tests, training, and exercises to ensure that agency personnel are familiar with assigned emergency roles and responsibilities and that the FDA EOP can be implemented rapidly and effectively.

Any deficiencies, findings, areas recommended for corrective action, or improvements arising from exercises will be addressed through additional training, plan updates, and/or demonstrations in subsequent emergency preparedness events.

G.6 IMPLEMENTATION

The FDA EOP, including the basic plan and all annexes, is effective for execution upon and pursuant to approval by the Commissioner.

OEM shall ensure this FDA EOP is subject to regular maintenance, review, and updates based on organizational and procedural changes, AARs, and new guidance. At any time, agency emergency coordinators, upon approval by the Center/Office senior official, should recommend to OEM improvements and changes thereto that are appropriate. The FDA EOP and any approved changes will be forwarded to all organizational components with responsibilities for implementation of the plan.

The FDA EOP shall be placed on the FDA websites (intranet) inside.fda and (Internet) www.fda.gov and updated as changes occur.
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Appendix A: FDA Support for the Emergency Support Function Annexes to the National Response Framework

The Emergency Support Function (ESF) annexes to the National Response Framework (NRF) provide the structure for coordinating Federal interagency support for a Federal response to an incident. These are mechanisms for grouping functions most frequently used to provide Federal support to States and Federal-to-Federal support, both for declared disasters and emergencies under the Robert T. Stafford Disaster Relief and Emergency Assistance Act (Stafford Act) and for non-Stafford Act incidents.

The ESF annexes are used by both Federal and State governments as the primary mechanism at the operational level to organize and provide assistance. ESFs align categories of resources and provide strategic objectives for their use. ESFs use standardized resource management concepts such as typing, inventorying, and tracking to facilitate the dispatch, deployment, and recovery of resources before, during, and after an incident.

Depending on the nature of the emergency, FDA’s emergency response roles and responsibilities may fall under one or more of the following ESFs.

A.1 ESF #8 – Public Health and Medical Services

The ESF #8 – Public Health and Medical Services Annex\(^4\) provides the mechanism for Federal assistance to supplement State, local, tribal, and territorial (SLTT) and insular area resources in response to a disaster, emergency, or incident that may lead to a public health, medical, behavioral, or human service emergency, including those that have international implications.

**Primary Agency**

- U.S. Department of Health and Human Services (HHS)

**ESF Coordinator**

- HHS

**Scope**

ESF #8 provides supplemental assistance to SLTT and insular area governments in the following core functional areas:

- Assessment of public health/medical needs
- Health surveillance
- Medical surge
- Health/medical/veterinary equipment and supplies
- Patient movement
- Patient care
- Safety and security of drugs, biologics, and medical devices
- Safety and security of blood and tissues
- Food safety and defense
- Agriculture safety and security
- All-hazards public health and medical consultation, technical assistance, and support
- Behavioral healthcare

\(^4\) For more information on ESF #8, refer to [www.fema.gov/media-library/assets/documents/32198?id=7359](http://www.fema.gov/media-library/assets/documents/32198?id=7359).
• Public health and medical information
• Vector control
• Guidance on potable water/wastewater and solid waste disposal
• Mass fatality management, victim identification, and decontaminating remains
• Veterinary medical support

**Core Capabilities and Actions**

The following table lists the response core capabilities that ESF #8 most directly supports along with the related ESF #8 actions.

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<tr>
<th>Table A-1. ESF Roles Aligned to Core Capabilities</th>
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<td>Core Capability</td>
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<td>Public Health Aspects of Potable Water/Wastewater and Solid Waste</td>
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Agency Actions
FDA will provide support, personnel, or expertise to HHS for many of the HHS responsibilities listed below. FDA will also:

- Monitor blood and tissue supplies, shortages, and reserves.
- Ensure the safety and security of food in coordination with other responsible Federal agencies (e.g., U.S. Department of Agriculture [USDA]). In cooperation with SLTT and insular area officials, assess whether food manufacturing, food processing, food distribution, food service, and food retail establishments in the affected area are able to provide safe food.
- In cooperation with SLTT and insular area officials as well as the food industry, conduct tracebacks and/or recalls of adulterated products.
- In cooperation with SLTT, insular area, and Federal officials, provide guidance regarding the proper disposal of contaminated products and the decontamination of affected food facilities in order to protect public health.
- Provide public health risk communication messages and advisories that communicate, in a culturally and linguistically appropriate and accessible manner, relevant information on health hazards or other situations that could potentially threaten public health.
- Disseminate public health information on protective actions related to exposure to health threats or environmental threats.
- Notify or respond to foreign country potential health threats as required by International Health Regulations (IHR).
- Provide support for public health matters for radiological incidents as a member of the Advisory Team for Environment, Food, and Health.
- Consult public health and medical subject matter experts (SMEs) with ESF #8 supporting organizations (including partners representing all appropriate populations such as pediatric populations, populations with disabilities and others with access and functional needs, the aging, and those with temporary or chronic medical conditions).

A.2 ESF #10 – Oil and Hazardous Materials Response
The ESF #10 – Oil and Hazardous Materials Response Annex provides Federal support in response to an actual or potential discharge and/or release of oil or hazardous materials (HazMat) when activated.

Primary Agencies
- U.S. Environmental Protection Agency (EPA)
- U.S. Department of Homeland Security (DHS)/U.S. Coast Guard (USCG)

ESF Coordinator
- EPA

Scope
ESF #10 may be activated as described in the NRF for a Stafford Act response, at the Secretary of Homeland Security’s discretion, and/or in response to a request for Federal-to-Federal support. Federal response to oil or HazMat incidents may also be carried out under another key Federal response authority.

45 For more information on ESF #10, refer to www.fema.gov/media-library/assets/documents/32241?id=7376.
called the National Oil and Hazardous Substances Pollution Contingency Plan (NCP), a regulation from 40 Code of Federal Regulations (CFR) Part 300. The NCP is an operational supplement to the NRF and may be used in conjunction with, or independent from, the Stafford Act.

The scope of ESF #10 includes the appropriate actions to prepare for and respond to a threat to public health, welfare, or the environment caused by actual or potential oil and HazMat incidents.

**HHS Actions**
HHS directly supports ESF #10 with the following action:

- **FDA**: Works in cooperation with EPA and USDA to ensure the proper disposal of contaminated food or animal feed.

### A.3 ESF #11 – Agriculture and Natural Resources

The ESF #11 – Agriculture and Natural Resources Annex organizes and coordinates Federal support for the protection of the Nation’s agricultural and natural and cultural resources during national emergencies. ESF #11 works during actual and potential incidents to provide nutrition assistance; respond to animal and agricultural health issues; provide technical expertise, coordination, and support of animal and agricultural emergency management; ensure the safety and defense of the Nation’s supply of meat, poultry, and processed egg products; and ensure the protection of natural and cultural resources and historic properties.

#### Primary Agencies
- USDA
- U.S. Department of the Interior (DOI)

#### ESF Coordinator
- USDA

#### HHS Functions
This section describes specific functions performed by HHS as an ESF #11 support agency:

- HHS National Veterinary Response Teams, veterinary officers of the Commissioned Corps of the U.S. Public Health Service (USPHS), and other veterinary medical professionals within its divisions (such as the Centers for Disease Control and Prevention [CDC], FDA, and the National Institutes of Health [NIH]) are the primary Federal resource for treatment of injured and/or ill pets, service animals, working animals, laboratory animals, and livestock. Under its own statutory authority, HHS can manage and conduct animal response to zoonotic diseases in order to protect human health.

- For livestock or poultry diseases exotic to the United States that are either not or only mildly zoonotic, HHS supports the USDA and its authority to manage a foreign animal disease response with the resources listed above.

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46 For more information on the NCP, refer to [www.epa.gov/OEM/content/lawsregs/ncpover.htm](http://www.epa.gov/OEM/content/lawsregs/ncpover.htm).

47 For more information on ESF #11, refer to [www.fema.gov/media-library/assets/documents/32213?id=7365](http://www.fema.gov/media-library/assets/documents/32213?id=7365).
Appendix B: FDA Support for the Incident Annexes to the NRF

The Incident Annexes to the NRF describe the concept of operations (CONOPS) to address specific contingency or hazard situations or an element of an incident requiring specialized application of the NRF.

Depending on the nature of the emergency, FDA’s response efforts support the following NRF Incident Annexes.

B.1 Biological Incident Annex

Coordinating Agency
- HHS

Purpose/Scope
The NRF Biological Incident Annex\(^\text{48}\) outlines the actions, roles, and responsibilities associated with a response to a human disease outbreak of known or unknown origin requiring Federal assistance. In this annex, a biological incident includes naturally occurring biological diseases (communicable and noncommunicable) in humans as well as terrorist events. This definition also includes those biological agents found in the environment, or diagnosed in animals, that have the potential for transmission to humans (zoonosis). The annex outlines biological incident response actions, including threat assessment notification procedures, laboratory testing, joint investigative/response procedures, and activities related to recovery.

The annex supports policies and procedures outlined in ESF #8, ESF #10, ESF #11, ESF #15 – External Affairs, the Terrorism Incident Law Enforcement and Investigation Annex, and the International Coordination Support Annex.

HHS serves as the Federal government’s primary agency for the public health and medical preparation and planning for and response to a biological terrorism or outbreak that results from either a known or novel pathogen, including an emerging infectious disease.

The USDA serves as the government’s primary agency for outbreaks and/or attacks that may occur in animals used in the commercial production of food. USDA may also serve as the government’s primary agency for attacks on food processing, including slaughtering facilities under its regulatory purview. In the event of a food or animal event, HHS may provide additional public health and veterinary epidemiological assistance to USDA. Wildlife events will be placed under the purview of DOI, while those involving marine animals will be managed and monitored by the U.S. Department of Commerce.

HHS Responsibilities
See ESF #8 for additional information.

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B.2 **Catastrophic Incident Annex**

**Coordinating Agency**
- DHS/Federal Emergency Management Agency (FEMA)

**Purpose/Scope**
The *Catastrophic Incident Annex* to the NRF (NRF-CIA)\(^{49}\) establishes the context and overarching strategy for implementing and coordinating an accelerated, proactive national response to a catastrophic incident. A catastrophic incident, as defined by the NRF, is any natural or man-made incident, including terrorism, that results in extraordinary levels of mass casualties, damage, or disruption severely affecting the population, infrastructure, environment, economy, national morale, and/or government functions.

During a catastrophic incident, normal procedures for certain ESFs may be expedited or streamlined to address the magnitude of urgent requirements of the incident. All ESFs must explore economies of scale to maximize utilization and efficiency of limited resources.

A more detailed and operationally specific *Catastrophic Incident Supplement* to the NRF (NRF-CIS) is published independently of the NRF and its annexes.

**FDA Responsibilities**
Upon notification from the National Operations Center (NOC) that the NRF-CIA has been implemented, Federal departments and agencies immediately:
- Take actions to activate, mobilize, and deploy incident-specific resources in accordance with the NRF-CIS.
- Take actions to protect life, property, and critical infrastructure under their jurisdiction and provide assistance within the affected area.
- Commence those hazard-specific activities established under the appropriate and applicable NRF Incident Annex(es), including the NRF-CIA.
- Commence functional activities and responsibilities established under the NRF ESF annexes.

All Federal departments and agencies and organizations assigned primary or supporting ESF responsibilities immediately begin implementation of those responsibilities as appropriate or when directed by the President.

B.3 **Food and Agriculture Incident Annex**

**Coordinating Agencies**
- USDA
- HHS

**Purpose/Scope**
The NRF *Food and Agriculture Incident Annex*\(^{50}\) describes the roles and responsibilities associated with all incidents involving the Nation’s agriculture and food systems that require a coordinated Federal response. The following are objectives of a coordinated national response to an incident impacting food and agriculture:

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• Detect the event through the reporting of illness, disease/pest surveillance, routine testing, consumer complaints, and/or environmental monitoring.

• Determine the primary coordinating agency.

• Determine the source of the incident or outbreak.

• Control and contain the distribution of the affected source.

• Identify and protect the population at risk.

• Assess public health, food, agriculture, and law enforcement implications.

• Assess the extent of residual biological, chemical, or radiological contamination, then decontaminate and dispose as necessary.

The NRF Food and Agriculture Incident Annex supports policies and procedures outlined in the NRF, ESF #8 – Public Health and Medical Services Annex, ESF #10 – Oil and Hazardous Materials Response Annex, ESF #11 – Agriculture and Natural Resources Annex, Terrorism Incident Law Enforcement and Investigation Annex, and the Federal Food and Agriculture Decontamination and Disposal Roles and Responsibilities document.

**HHS Responsibilities**
Specific responsibilities are described in greater detail in ESF #8, ESF #10, and ESF #11.

HHS provides leadership by ensuring the safety and security of food, animal feed, food-producing animals, and animal therapeutics. HHS, through the CDC and in coordination with States, develops and implements surveillance systems to monitor the health of the human population.

HHS, through FDA, has statutory authority for all domestic and imported food except meat, poultry, and egg products that are under the authority of the USDA Food Safety and Inspection Service (FSIS). FDA also has statutory authority for animal feed and for the approval of animal drugs intended for both therapeutic and nontherapeutic use in food animals as well as household pets and service animals.

**FDA Responsibilities**
FDA responsibilities are as follows:

• If FDA suspects a threat involving biological or chemical agents, nuclear/ radiological material or indications that instances of disease may not be the result of natural causes, the U.S. Department of Justice (DOJ) must be notified through the DOJ Federal Bureau of Investigation Weapons of Mass Destruction (FBI WMD) Operations Unit.

• Any potential or actual incidents requiring a coordinated Federal response involving contaminated food, infected animals or plants, or economically devastating plant pest infestation should be reported to the Secretary of Homeland Security through the NOC at 202-282-8101 and brought to the attention of designated officials according to ESF #8 and ESF #11.

• If terrorist activity is suspected in connection with the incident, procedures outlined in the Terrorism Incident Law Enforcement and Investigation Annex will be followed.

**B.4 NUCLEAR/RADIOLOGICAL INCIDENT ANNEX**

**Coordinating Agencies**

• U.S. Department of Defense (DoD)
• U.S. Department of Energy (DOE)
• DHS
Purpose/Scope

The Nuclear/Radiological Incident Annex (NRIA) to the NRF describes the policies, situations, CONOPS, and responsibilities of the Federal departments and agencies governing the immediate response and short-term recovery activities for incidents involving release of radioactive materials to address the consequences of the event. These incidents may occur on federally owned or licensed facilities, privately owned property, urban centers, or other areas and may vary in severity from the small to the catastrophic. The incidents may result from inadvertent or deliberate acts. The NRIA applies to incidents when the nature and scope of the incident requires a Federal response to supplement the State, tribal, or local incident response.

This annex applies to two categories of nuclear and radiological incidents: (1) inadvertent or otherwise accidental releases and (2) releases related to deliberate acts. These incidents may also include potential release of radioactive material that poses an actual or perceived hazard to public health, safety, national security, and/or the environment. The category covering inadvertent releases includes: two categories of nuclear facilities (commercial or weapons production facilities), lost radioactive material sources, transportation accidents involving nuclear/radioactive material, domestic nuclear weapons accidents, and foreign accidents involving nuclear or radioactive material that impact the United States or its territories, possessions, or territorial waters. The second category includes, but is not limited to, response to the effects of deliberate attacks perpetrated with radiological dispersal devices (RDDs), nuclear weapons, or improvised nuclear devices (INDs).

This annex applies whenever a Federal response is undertaken unilaterally pursuant to Federal authorities or when an incident exceeds or is anticipated to exceed State, tribal, or local resources. The level of Federal response to a specific incident is based on numerous factors, including the ability of State, tribal, and local officials to respond; the type, amount, and custody of (or authority over) radioactive material involved; the extent of the impact or potential impact on the public and environment; and the size of the affected area.

FDA Responsibilities

FDA responsibilities are as follows:

- Provide a representative to the Advisory Team to develop coordinated advice and recommendations on environmental, food, health, and animal health matters for the Incident Command/Unified Command (UC), DHS, Joint Field Office (JFO) Unified Coordination Group, the coordinating agency, and/or State, tribal, and local government as appropriate.

- Provide laboratory capabilities for food and agriculture analysis.
  - FDA must maintain laboratory capability for regulatory, confirmatory, and screening analysis for both routine and emergency response operations including:
    - Maintaining an alternate laboratory location for both routine and emergency response confirmatory analysis in the case of an emergency shutdown of the Winchester Laboratory
    - FDA must maintain Food Emergency Response Network (FERN) capability to screen and triage food and agricultural samples for response to a radiological emergency.
B.5 TERRORISM INCIDENT LAW ENFORCEMENT AND INVESTIGATION ANNEX

Coordinating Agency

- DOJ/FBI

Purpose/Scope

The purpose of the Terrorism Incident Law Enforcement and Investigation Annex is to facilitate an effective Federal law enforcement and investigative response to all threats or acts of terrorism within the United States, regardless of whether they are deemed credible and/or whether they escalate to an “incident of national significance.” To accomplish this, the annex establishes a structure for a systematic, coordinated, unified, timely, and effective national law enforcement and investigative response to threats or acts of terrorism within the United States.

This annex is a strategic document that:

- Provides planning guidance and outlines operational concepts for the Federal law enforcement and investigative response to a threat or an actual terrorist incident within the United States.
- Acknowledges and outlines the unique nature of each threat or incident, the capabilities and responsibilities of the local jurisdictions, and the law enforcement and investigative activities necessary to prevent or mitigate a specific threat or incident.

FDA Responsibilities

FDA responsibilities are as follows:

- Provide support to Federal law enforcement and investigative response efforts related to threats or acts of terrorism involving or impacting FDA-regulated products.
- If terrorist activity is suspected in connection with any incident involving FDA-regulated product, the Terrorism Incident Law Enforcement and Investigation Annex will be followed and the Office of Criminal Investigations (OCI) should be notified of such. OCI will coordinate the terrorism incident law enforcement response with the FBI.
- Provide laboratory capabilities for food and agriculture analysis.
  - FDA must maintain essential laboratory capability for testing of its regulated products.
  - ORA laboratory network provides redundant capability in key areas such that analytical operations associated with an emergency can still be conducted in the event of a shutdown of one of the laboratories.
  - The FDA FERN includes a group of labs supported by the agency Cooperative Agreement Program (CAP) that have key microbiological, chemical, and radiological testing capability. These CAP labs can also be utilized in an emergency situation to provide essential testing in FDA priority areas as well.
Appendix C:
Processing Mission Assignment Sub-Taskings

C.1 PURPOSE

The purpose of this appendix is to establish a uniform procedure for FDA to execute sub-tasks under mission assignments (MAs). The appendix supports HHS MAs under the NRF and the National Disaster Recovery Framework (NDRF).

FDA provides timely disaster response operating under its own authorities. In response to a catastrophic event, there are cases where DHS/FEMA will issue MAs to HHS/FDA to provide additional resources to States and local jurisdictions affected by the disaster. Types of assignments include supporting States for the inspection of retail food establishments and pharmacies, sample collection, and providing SMEs. A list of possible assignments is provided at Attachment C-1 (FDA Pre-Scripted Mission Assignments).

MAs are addressed in the FDA Emergency Operations Plan (EOP) in Section B.2.2.3.4, “Mission Assignments in Support of State and Local Response.”

C.2 ASSUMPTIONS

Assumptions are as follows:

- MAs may be issued when the NRF or NDRF is activated as part of a Stafford Act declaration that requires a coordinated Federal response to manage a disaster and/or emergency. MAs may also be issued to HHS when Recovery Support Function (RSF) annexes are activated under the NDRF.
- The need for Federal assistance exists when the incident is of such severity and magnitude that SLTT resources are exceeded. The SLTT requests support activities outside the normal scope of FDA’s mission.
- This appendix refers only to MAs issued by FEMA to HHS and not requests for assistance from other Federal and/or SLTT agencies. The term “mission assignment” may at times be used to describe a variety of missions and tasks undertaken by FDA.
- FDA completes assignments only as MA sub-tasks issued by HHS. FDA does not sub-task other Federal agencies. Offers of assistance from other Federal agencies should be referred to the HHS SOC or FEMA.
- MA funds are used appropriately and expeditiously. Financial operations are conducted in accordance with established Federal law, policies, regulations, and standards to protect Federal government resources from waste, fraud, and mismanagement.

C.3 SCOPE

This appendix applies to all FDA personnel involved in processing an HHS MA sub-task, including FDA field and headquarters staff and HHS response elements including the EMG Operations, the Human Services Watch Officer, SOC Operations Officers, and ASPR Administration and Finance (A&F).
C.4 Procedure

C.4.1 Processing Resource Request Form

C.4.1.1 FDA/State Coordination

While the State is responsible for determining their specific needs during a disaster, FDA field staff may assist the State in developing language for inclusion into Resource Request Forms (RRFs) if requested (see Attachment C-2 [RRF for FDA Retail Food Establishment Inspections Sample – FEMA Form 010-0-7]). The MA sub-task process allows FDA to be reimbursed for providing States with additional resources and assistance beyond the services and guidance provided by FDA during normal operations. Figure C-1 provides an overview of the process.

The MA sub-task process is as follows:

1. Information exchanges with the State can be initiated by District offices on an informal basis to ensure requests are within FDA’s capabilities and to facilitate coordination. To expedite this step, see Attachment C-1 for a list of FDA Pre-Scripted Mission Assignments (PSMAs) prepared specifically for FDA as HHS MA sub-tasks. PSMAs provide examples and templates for the State to use in developing specific requests and submitting them in RRFs. See Attachment C-2 for instructions on completing RRFs and a sample RRF.

2. Once FDA field begins discussions above with the State, the Office of Emergency Management/Office of Emergency Operations (OEM/OEO) should be notified of a possible RRF to be submitted to FEMA.

3. During the information exchange, it may be necessary to assist the State to define requests. Specific State requirements should be identified and compared with State capabilities. If a gap is identified, appropriate FDA resources should be identified for meeting shortfalls in coordination with OEM/OEO.
4. The State is responsible for initiating, reviewing, and submitting all RRFs to FEMA for review, approval, and submission to the appropriate department/agency/ESF. Attachment C-1 also provides a draft statement of work developed by FDA.

5. HHS may deploy an Incident Response Coordination Team (IRCT) to be primarily responsible for supporting the Federal public health and medical management of an incident. The IRCT provides the field management component of the Federal public health and medical response. FDA field staff will advise the IRCT as appropriate regarding the scope of any RRF. FDA may provide a liaison to the IRCT.

6. FEMA receives and reviews the RRF to determine if the request is appropriate for the specific event. If FEMA agrees, the RRF is submitted to the appropriate ESF agency for their review, cost estimation, and return to FEMA for final review and approval.

**C.4.1.2 Initial FDA Internal Coordination (Field and Headquarters)**

Initial FDA internal coordination (field and headquarters) including the following:

1. Activation of an ESF/RSF, where FDA is a support agency, may indicate that MAs may be sub-tasked to FDA. For incidents that may require FDA support to the States, OEM/OEO will inform the appropriate FDA Centers/Offices that activation has occurred and MA sub-tasks may be received.

2. If an FDA Incident Management Group (IMG) is established in the FDA Emergency Operations Center (EOC), MA sub-task reviews will be performed by the appropriate IMG member. If an IMG is not established, MA sub-task reviews will be performed by OEM/OEO. From this point forward in the appendix, the term “FDA EOC” refers to the IMG or OEM/OEO.

3. If an Incident Management Team (IMT) is established by an FDA District Office, MA sub-task reviews will be performed by the appropriate IMT member designated by the Incident Commander (IC). If an IMT is not established, MA sub-task reviews will be performed by the appropriate staff from the Program or District Office.

**C.4.2 Processing a Mission Assignment Sub-Task**

FEMA may issue MAs to other Federal agencies to (1) address a State’s request for Federal assistance to meet emergency needs or (2) support overall Federal operations pursuant to, or in anticipation of, a Stafford Act declaration.

**C.4.2.1 FDA/FEMA/HHS Coordination**

FDA/FEMA/HHS coordination will be as follows:

1. The FEMA JFO will develop an MA (see Attachment C-3 [MA Form Based on Completed RRF Sample – FEMA Form 010-0-8]) based on the RRF. MA development will take place with input and review by the District Director (DD), or designee from the relevant District, in consultation with the FDA EOC, and in coordination with the HHS Regional Emergency Coordinator (REC). The cost associated with the execution of the MA will be based on PSMA cost estimates or other cost information from the FDA Office of Regulatory Affairs (ORA) and/or relevant Centers.

2. FEMA will task HHS by sending the formal MA with the FEMA Integrated Financial Management Information System (IFMIS) stamp (certification of funds) to the HHS SOC (hhs.soc@hhs.gov).

3. The FDA EOC will alert the appropriate Centers/Offices that an MA sub-task is forthcoming.
4. Upon approval, HHS/ASPR A&F will send the MA to HHS EMG Operations for the final approval before sub-tasking FDA.

5. The HHS/ASPR A&F will send the signed sub-task to the FDA EOC obligating funds for FDA to respond to the State’s request.

6. The FDA EOC will forward the sub-task to the ORA Office of Resource Management (ORM) and Office of Financial Management (OFM) for review and signature. Based on their review, ORA ORM/OFM will provide final review and signature and forward back to the FDA EOC.

7. The FDA EOC will scan the signed sub-task, enter it into the Emergency Operations Network – Incident Management System (EON-IMS), and send it back to the HHS SOC with a copy to HHS/ASPR A&F.

C.4.2.2 FDA Internal Coordination

FDA internal coordination is as follows:

1. The FDA EOC notifies appropriate Program and District staff and Centers/Offices of the sub-task approval.

2. Sub-tasks will be documented in the Incident Action Plan (IAP).

3. The FDA EOC transmits the sub-task funding requirements to the Office of the Commissioner (OC)/Office of Operations (OO)/Office of Finance, Budget, and Acquisition (OFBA)/Office of Financial Operations (OFO)/OFM/Division of Budget Execution and Control.

4. The Division of Budget Execution and Control informs the Office of Global Regulatory Operations and Policy (OGROP)/ORA/ORM/Division of Budget Formulation and Execution, which is responsible for issuing accounting data and providing funding to the District Office for response actions. OGROP/ORA/Office of Policy and Risk Management (OPRM)/Division of Planning, Evaluation, and Management may establish a Program Account Code (PAC) or refer to an existing one to provide for time reporting. Response personnel enter data (such as laboratory and investigation time expended) into the Field Accomplishments and Compliance Tracking System (FACTS) on an ongoing basis.

5. FDA field staff for the affected area notifies the FEMA JFO or District Offices, the affected State(s), and others as appropriate of the receipt of a valid sub-tasking from HHS and corresponding authorization to take appropriate response actions.

C.4.2.3 MA Monitoring and Closeout

The MA monitoring and closeout process is as follows:

1. During operations under the sub-task, the FDA EOC will monitor the expenditure of funds in order to avoid potential over-spending. When the obligation level approaches 75 percent of authorized funds, the FDA EOC will notify HHS/ASPR A&F if any additional funding needs are anticipated. HHS/ASPR A&F will coordinate with FEMA to amend the MA as required. Justification for additional funding is required. The FDA EOC needs to be kept apprised of any changes in the scope of the sub-task.

2. When nearing completion of the mission, FDA field staff will discuss with the appropriate FEMA JFO ESF lead and State representatives any issues regarding the completion of the work and ensure the work has been completed in a satisfactory manner. Decisions regarding the official end date of the mission will be made during this communication.
3. Upon deactivation of the MA, FDA field staff must provide the FEMA MA Manager (with a courtesy copy to the Human Services Watch Officer) with a listing of any other ongoing sub-tasks and their status, Property Disposition Forms (if applicable), and financial and program points of contact (POCs).

4. Prior to demobilization, any deployed FDA personnel will notify their supervisor and the FDA EOC of travel arrangements.

5. The FDA EOC will compile and submit all data, costs, and supporting documentation as requested in the Stafford Act Mission Assignment Sub-Tasking Form (see example in Attachment C-4) to HHS/ASPR A&F.

6. HHS/ASPR A&F is responsible for submitting all close-out reports to the FEMA MA office for any additional eligible reimbursable costs incurred during the mission.
Requests for support from State and local agencies can be made directly to FDA or, in the case of a declaration of a major disaster or emergency, through FEMA. In order to assist States and local agencies with such requests, FDA has developed PSMAs under ESF #8, which can be used as examples to expedite the preparation and submission of such requests. Draft MA text is provided below.

<table>
<thead>
<tr>
<th>Assistance Requested</th>
<th>HHS will provide food and product safety investigations to augment State and local staff in support of disaster operations in response to [insert disaster name/#].</th>
</tr>
</thead>
<tbody>
<tr>
<td>Statement of Work</td>
<td>At the direction of, and in coordination with FEMA, HHS will address immediate and short-term issues that impact whether food, drugs, biologics, and medical devices that have been directly affected by this incident are still appropriate for use. HHS will also assist in determining what steps, if any, may be employed to restore drugs, biologics, medical devices, and food to a condition whereby they would be fit for use. FDA investigators and consumer safety officers will assist State and local staff with the following services:</td>
</tr>
<tr>
<td></td>
<td>• Collect and analyze samples of foods, drugs, cosmetics, and/or medical devices for attributes, as necessary, to assist in providing assurance that commodities are fit for use.</td>
</tr>
<tr>
<td></td>
<td>• Conduct food safety inspections at retail food service establishments to ensure compliance with food safety standards, to include restaurants, school and hospital cafeterias, daycare center food service establishments, temporary shelters, and others.</td>
</tr>
<tr>
<td></td>
<td>• Conduct biologic, drug, and medical product safety inspections of pharmacies, hospitals, retail establishments, and other establishments that provide human and/or animal drugs, biologics, and medical devices to ensure these items have been stored under appropriate conditions and are fit for use.</td>
</tr>
<tr>
<td></td>
<td>• Conduct assessments for acceptable levels of radiation emission in facilities where diagnostic x-ray and mammography equipment are installed.</td>
</tr>
<tr>
<td></td>
<td>• Train disaster response personnel in food safety, preparation, handling, and storage.</td>
</tr>
</tbody>
</table>

| Lease or Purchase of Accountable Property | None. |

| FEMA Logistical Support | • Teams may require some level of base operating support (e.g., food, shelter, laundry). |
|                        | • MA task orders will be issued for individual personnel and/or teams as required by the incident and will specify times, location, and dates. |
|                        | • HHS may sub-task other Federal agencies for disaster medical support requirements as needed. |
Type of Assistance
Direct Federal assistance.

Cost depends on which of the following resources are utilized. All teams will not necessarily be deployed:

- **FDA Retail Food Establishment Inspections.** FDA investigators (consumer safety officers) to augment State and local staff to perform inspections of establishments serving food at retail in <location> for conformance to appropriate food safety standards. Such establishments may include restaurants, school and hospital cafeterias, daycare center food service establishments, and temporary shelters, among others.

- **FDA Food, Cosmetic, and Medical Product Establishment Inspection.** FDA investigators (consumer safety officers) to augment State and local staff to conduct inspections of establishments that prepare, pack, and/or hold human and/or animal food, human and/or animal drugs, biologics, cosmetics, and/or medical devices to help ensure such commodities are safe, effective, and/or otherwise fit for use.

- **FDA Pharmacy Inspections.** FDA investigators (consumer safety officers) to augment State and local staff to perform inspections of pharmacies and other establishments offering human and/or animal drugs, biologics, and medical devices at retail to assist in ensuring such drugs, biologics, and medical devices have been stored under appropriate conditions and are fit for use.

- **FDA Sample Collections.** FDA investigators (consumer safety officers) to augment State and local staff to perform sample collections of human and/or animal foods, cosmetics, human and/or animal drugs, biologics, and medical devices for subsequent analyses.

- **FDA Analysts.** FDA analysts (including microbiologists, chemists, biologists, and other disciplines as appropriate) to augment State and local staff to analyze samples of foods, drugs, medical devices, and cosmetics to assist in providing assurance that these commodities are fit for use.

- **FDA Food Safety Training.** FDA investigators (consumer safety officers) or other FDA staff with food safety expertise to provide training in food safety preparation, handling, and storage to volunteers and/or other appropriate disaster response personnel.

- **FDA SMEs.** FDA SMEs to augment State and local staff to address issues that impact whether human and/or animal drugs, biologics, human and/or animal foods, and medical devices are appropriate for use and/or to provide guidance on what steps, if any, may be employed to restore human and/or animal drugs, biologics, human and/or animal foods, cosmetics, and medical devices to a condition whereby they would be fit for use.

- **FDA X-Ray Mammography Analysis.** FDA investigators (consumer safety officers) with expertise in this area to augment State and local staff to conduct assessments (field tests) of facilities where diagnostic x-ray and mammography equipment are installed, to help ensure the equipment is operating within acceptable radiation emission limits.
Attachment C-2: RRF for FDA Retail Food Establishment Inspections Sample

Instructions on Completing the Resource Request Form (RRF)

The State requesting assistance will need to complete Section I and Section II; most items are self-explanatory. For item II.1 (Description of Requested Assistance), the appropriate PSMA can form the basis for the text to be inserted. PSMAs serve only as a guide to expedite the process; the text can be modified as needed to reflect the mission, local conditions, and State requirements.

In addition, HHS requires a preliminary estimate of costs based upon the number of resources for the duration deployment. Use the following as a basis for an estimate:

1. **Deploy one investigator for 7 days**: $5,000
   (does not include standard pay; includes travel, per diem, and overtime)

   Calculate:
   
   \[ \text{Weeks} \times \text{Personnel Deployed} \times 5,000 = \text{Estimated Cost} \]

   **Example**:
   
   2 weeks x 5 personnel x 5,000 = $50,000

   **NOTE**: The costs are only estimates that should be used for development of RRFs. These estimates may be adjusted by HHS/ASPR A&F or FEMA prior to issuance of the MA. If the funding amount on the sub-task is below the estimated cost provided by FDA, the Program and/or District Office should notify the FDA EOC, who will discuss with HHS/ASPR A&F.
**RESOURCE REQUEST FORM (RRF)**

**PAPERWORK BURDEN DISCLOSURE NOTICE**

Public reporting burden for this form is estimated to average 30 minutes per response. The burden estimate includes the time for reviewing instructions, searching existing data sources, gathering and maintaining the needed data, and completing and submitting this form. This collection of information is required to obtain or retain benefits. You are not required to respond to this collection of information unless it displays a valid OMB control number. Send comments regarding the accuracy of the burden estimate and any suggestions for reducing this burden to: Information Collections Management, Department of Homeland Security, Federal Emergency Management Agency, 500 C Street, SW, Washington, DC 20472-3100, Paperwork Reduction Project (1660-0047). NOTE: Do not send your completed form to this address.

I. REQUESTING ASSISTANCE (To be completed by Requestor)

1. Requestor’s Name (Please print): Kurt Donaldson
2. Title: Program Manager
3. Phone No.: 806-555-6566
4. Requestor’s Organization: CMOH
5. Fax No.
6. E-Mail Address

II. REQUESTING ASSISTANCE (To be completed by Requestor)

1. Description of Requested Assistance:

2. Quantity
3. Priority [ ] Lifesaving [ ] Life Sustaining [X] High
4. Date and Time Needed

5. Delivery Site Location
6. Site Point of Contact (POC):
   - Telephone No.
   - Fax No.

9. State Approving Official Signature
10. Date and Time

III. SOURCING THE REQUEST - REVIEW AND SIGNATURE (Operations Section Only)

<table>
<thead>
<tr>
<th>1.</th>
<th>Source:</th>
</tr>
</thead>
<tbody>
<tr>
<td>[X] OPS Review by:</td>
<td></td>
</tr>
<tr>
<td>[ ] LOG Review by:</td>
<td></td>
</tr>
<tr>
<td>Other Coordination:</td>
<td></td>
</tr>
<tr>
<td>[ ] Other Coordination:</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>2.</th>
<th>Assigned to:</th>
</tr>
</thead>
<tbody>
<tr>
<td>[X] Donations</td>
<td></td>
</tr>
<tr>
<td>[ ] Other (Explain)</td>
<td></td>
</tr>
<tr>
<td>[ ] Requisitions</td>
<td></td>
</tr>
<tr>
<td>[ ] Procurement</td>
<td></td>
</tr>
<tr>
<td>[ ] Interagency Agreement</td>
<td></td>
</tr>
<tr>
<td>[ ] Mission Assignment</td>
<td></td>
</tr>
</tbody>
</table>

| 3. | Date/Time: |

4. Immediate Action Required [ ] Yes [ ] No

IV. STATEMENT OF WORK (Operations Section Only)

1. OFA Action Officer
2. 24 Hour Phone #
3. Fax #

4. FEMA Project Manager
5. 24 Hour Phone #
6. Fax #

7. Statement of Work [ ] See Attached

8. Estimated Completion Date
9. Estimated Cost

V. ACTION TAKEN (Operations Section Only)

| [ ] Accepted |
| [ ] Rejected |
| [ ] Requestor Notified |

Reason / Disposition

FEMA FORM 010-0-7
PREVIOUSLY FF 90-135
Page 1 of 2

**Attachment C-2: RRF for FDA**

**Retail Food Establishment Inspections**

July 2019
RESOURCE REQUEST FORM (RRF)

**TRACKING INFORMATION** (FEMA Use Only)

<table>
<thead>
<tr>
<th>ECAPS/NEMIS Task ID:</th>
<th>Resource Request #</th>
<th>Program Code/Event #</th>
<th>Originated as verbal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Received by (Name and Organization)</td>
<td>State</td>
<td>Date/Time Received</td>
<td></td>
</tr>
</tbody>
</table>

**INSTRUCTIONS**

Items on the Resource Request form that are not specifically listed are self-explanatory. Indicate “see attached” in any field for which additional space or more information is required.

I. Who is requesting assistance? Completed by requestor.

II. What needs to be done? Completed by requestor.

**Description of Requested Assistance:** Detail of resource shortfalls, statement of deliverable, or simply state problem/need.

**Priority:** The requestor’s priority, which may differ from the priority in BOX III.

**Site PC:** The person at the delivery site coordinating reception and utilization of the request/assistance. 24-hour contact information required.

If for Direct Federal Assistance (DFA), State Approving Official: Signature certified.

(1) State and local governments cannot perform, nor can they control, the performance of the requested work;

(2) Work is required as a result of the event, not a state action, such as demand for services;

(3) The State is providing the required assurances as set forth in 44 CFR, Section 206.

III. Action Review/Coordination (OPS Section Use Only): Completed by Operations Section Chief or Resource Capability Branch Director.

**Accept/Reject:** Operations Section Chief or Resource Capability Branch Director accepts or rejects the request; provide reason if rejection. If request accepted, coordinating with others (i.e., other Directors or Group Supervisors, begins to determine means of fulfilling request. All involved in coordination should check acceptance and initial or print their name.

**Assigned to:** Operations Section Chief or Resource Capability Branch Director assigns tasks. Orignation, may indicate the OPA Action Officer. Operations Section Chief may reassign the Action Officer if known, or tasked organization may make this assignment. This may be Emergency Support Function of the FEMA Organization (i.e., Logistics), or other organization.

**Date/Time Assigned:** Operations Section Chief or Resource Capability Branch Director provides date and time of when sourcing should begin.

IV. Statement of Work (OPS Section Use Only): Completed by the Operations Section Chief or Resource Capability Branch Director.

**OFA Action Officer:** Ops Section Chief obtains from OFA if request fulfilled by a MA; 24-hr phone/fax required. Information used in eCAPS.

**FEMA Project Manager:** Provided by Operations Section Chief, a Region PFT; 24-hr phone/fax required. Information used in eCAPS.

**Statement of Work:** Description of tasks to be performed. Could be to assess a problem and report back, or could be to proceed with a specific action. If 40-1 or MA, this goes in “justification” tab in eCAPS.

V. Action Taken (OPS Section Use Only): Completed by Operations Section Chief, Resource Capability Branch Director. MA Unit or Logistics.

**Resource Request Results:** Ops Section Chief, Resource Support Section Chief, MA Unit, or LOG should note what type of document the action resulted in by “checking” the appropriate box i.e., Mutual Aid, Donations, Requisition, Procurement, IA, MA, Other. If “Other” is selected, write in appropriate response or state “see below” and give detail description in “Disposition” field. “Disposition” field should note steps taken to complete the Action, and personnel, sub-tasked agencies, contacts and other resources utilized.

**TRACKING INFORMATION:** Completed by Action Tracker. Required for all requests.
Attachment C-3: MA Form Based on Completed RRF Sample

DEPARTMENT OF HOMELAND SECURITY
Federal Emergency Management Agency
MISSION ASSIGNMENT (MA)

PAPERWORK BURDEN DISCLOSURE NOTICE
Public reporting burden for this form is estimated to average 20 minutes per response. The burden estimate includes the time for reviewing instructions, searching existing data sources, gathering and maintaining the needed data, and completing and submitting this form. This collection of information is required to obtain or retain benefits. You are not required to respond to this collection of information unless it displays a valid OMB control number. Send comments regarding the accuracy of the burden estimate and any suggestions for reducing this burden to: Information Collections Management, Department of Homeland Security, Federal Emergency Management Agency, 500 C Street, SW, Washington, DC 20772-3100, Paperwork Reduction Project (1660-0047). NOTE: Do not send your completed form to this address.

I. TRACKING INFORMATION (FEMA Use Only)
State
CM (State of Columbia) Incident: 2017102505
Resource Request Number: 1008-108971
Program Code/Event Number
4088R-CM Hurricane Zulu
Date/Time Received: 12/04/2017 23:00

II. ASSISTANCE REQUIRED
Assistance Requested
HHS will provide food and product safety investigators to augment state, tribal and local staff in support of disaster operations in response to Hurricane Zulu in the State of Columbia.

III. INITIAL FEDERAL COORDINATION
Action:
ESF # 6: Internal Control Number: RRF #160
Site POC Name: KARL DONALDO
Initiator/Requestor Name: CARL DONALDO
24 Hour Phone: (609) 555-9800
Email Address: karl.donaldo@state.com
Date: 12/04/2017
Data/Time Requested: 12/04/2017 22:52

IV. DESCRIPTION (Job Category/Location)

V. COORDINATION (FEMA Use Only)
Type of MA:
Direct Federal Assistance
State Cost Share Percent: 25%
State Cost Share Amount: $ 98,250

STATEMENT OF WORK
At the direction of and in coordination with FEMA, HHS will provide food and product safety investigators to augment state and local staff to perform inspections of establishments serving food at retail establishments for conformance to appropriate food safety standards. Such establishments may include restaurants, school and hospital cafeterias, day care centers, foster care facilities, temporary shelters, etc.

Assignee Agency:
HHS (DEPT. OF HEALTH & HUMAN SVC.)
Project/Start Date: 12/09/2017
Estimated Projected End Date: 1/31/2018

New or Amendment to MA #: 
Total Cost Estimated: $35,000.00
Total Required this Obligation Cycle:

ESF/OFAR/ARSF Action Officer:
RONALD LENZI
Phone #: (305) 555-0400
Email: ronald.lenzi@hhs.gov

Mission Assignment Manager (Preparer):
MINDY MCDONALD
Date: 12/05/2017

*FEMA Project Manager/Branch Director (Program Approval)
JAMES DONALDO
Date: 12/05/2017

**Controller/Funds Control (Funds Review)
KENNY JEFFERS
Date: 12/05/2017

PREVIOUSLY FF 90-126 Page 1 of 2
### MISSION ASSIGNMENT (MA)

#### VI. APPROVAL

<table>
<thead>
<tr>
<th>*State Approving Official (Required for DFA)</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Federal Approving Official (Required for all)</strong></td>
<td>Date</td>
</tr>
<tr>
<td>JACK GRAHAM</td>
<td>12/05/2017</td>
</tr>
</tbody>
</table>

#### VII. OBLIGATION (FEMA Use Only)

<table>
<thead>
<tr>
<th>Mission Assignment Number</th>
<th>Amount This Action</th>
<th>Date/Time Obligated</th>
</tr>
</thead>
<tbody>
<tr>
<td>4088</td>
<td>$365,000</td>
<td>12/06/2017</td>
</tr>
</tbody>
</table>

**Signature required for all MAs.**

---

**INSTRUCTIONS**

Items on the Mission Assignment (MA) form that are not listed are self-explanatory.

**I. TRACKING INFORMATION.** Completed by Resource Support Section or Operations staff. Required for all requests.

**State:** If multi-State, choose State most likely to receive resource(s), i.e., when using 7220-SU Program Code

**Resource Request No.:** Based on chronological log number Used for tracking.

**Program Code/Event No.:** The pre-declaration, emergency, or major disaster number assigned for funding the event. Examples: 7220-SU, 3130-EM, 1248-DR.

**II. ASSISTANCE REQUESTED.** Completed by requestor.

**Assistance Requested:** Details from the Resource Request Form will provide for an action concerning assistance requested.

**Initiator/Requestor:** The initiator may be an individual filling the position of COE or a person signing a request on behalf of the POC.

**POC Name:** The person coordinating reception and utilization of requested resources and 24-hour contact information required.

**III. INITIAL FEDERAL COORDINATION.** Completed by FEMA personnel with Delegated Authority.

**Action to:** May be Emergency Support Function (ESF), Support Function (SF), internal FEMA organization, or other organization, which assigns the Action Officer.

**Remainder of MA used only to identify Federal agency to perform reimbursable work under (MA). Deliberate validation and verification of information must occur. MA is completed and issued.**

**IV. DESCRIPTION.** Completed by assigned agency Action Officer.

**Statement of Work:** Detailed description of work to be performed that includes: Overview of MA, objectives, tasks, resources, personnel, deliverable, location, period of performance and comprehensive cost estimate for period of performance. Statement of Work may be attached.

**Additional guidance concerning the writing of a Statement of Work can be found in the Mission Assignment Guide and PAR.**

**Assigned Agency:** Agency receiving the MA from FEMA. Activities within the scope of an ESF/RSF result in an MA to primary agency. Cite subordinate organization if applicable. Example: DOT-FAA, COE-SAD.

**Projected Start/End Date:** If end date is not clear, estimate and budget for 30, 60, or 90 days, then reevaluate. TBO is not acceptable; a date must be entered.

**Total Cost Estimate:** Enter dollar value and attach a detailed budget outlining personnel, equipment, contract, sub-tasked agency, travel and other costs. The cost estimate should include the total cost projection for the MA across the entire length of the MA. The 90 day obligation cycle is used to obligate funding in 90 day increments when completion period is expected to exceed 90 days.

**V. COORDINATION.** Completed by MA, except for Project Manager and Comptroller signatures.

**Type of MA:** Select only one.

**Appropriation Code:** Static data. Do not change. This is for information only, should not be used to report internal agency finances to Treasury.

**Receiving MA agencies are required to provide reporting as determined by the Program Manager.**

**VI. APPROVAL.** Completed by State Approving Official and Federal Approving Official.

**VII. OBLIGATION.** Completed by Financial Specialist.

**Mission Assignment No.:** Assigned in FEMA financial system chronologically using assigned agency acronym and two digit number.

**Amendment No.:** Note supplement number. For example: COE: SAD-01, Supp. 1, or DOR-08, Supp. 2.

**Amount this Action:** Taken from total cost estimate above.

**Cumulative Amount:** Cumulative amount for this MA, including amendments.
Attachment C-4: Stafford Act Mission Sub-Tasking Based on MA Form Sample

Stafford Act Mission Assignment Sub-Tasking Form

<table>
<thead>
<tr>
<th>Mission Assigned Primary Agency:</th>
<th>U.S. Department of Health and Human Services (HHS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sub-Tasked Agency:</td>
<td>U.S. Food and Drug Administration (FDA)</td>
</tr>
<tr>
<td>Tasking Statement/Statement of Work:</td>
<td>Provide food and product safety investigators to augment State, tribal, and local staff in support of disaster operations in response to Hurricane Zulu and in the State of Columbia. This tasking covers incremental salaries and travel expenses starting on December 6, 2017.</td>
</tr>
<tr>
<td>Project Completion Date:</td>
<td>January 31, 2018 (may be extended by email from Mr. David Dolinsky)</td>
</tr>
<tr>
<td>Authorized Funding:</td>
<td>$380,000</td>
</tr>
<tr>
<td>Accounting Information:</td>
<td>FDA shall bill and provide supporting documentation to HHS/ASPR using the following information:</td>
</tr>
<tr>
<td></td>
<td><strong>FL</strong></td>
</tr>
<tr>
<td>Agency Location Code (ALC):</td>
<td>75060099</td>
</tr>
<tr>
<td>Appropriation Number:</td>
<td>47-31-1004</td>
</tr>
<tr>
<td>Mission Assignment Number:</td>
<td>4086DR-CM-HHS05</td>
</tr>
<tr>
<td>CA Account Number:</td>
<td>1996389</td>
</tr>
<tr>
<td>Object Class:</td>
<td>Multiple</td>
</tr>
<tr>
<td>Amount This Action:</td>
<td>$380,000</td>
</tr>
<tr>
<td>DUNS Number:</td>
<td>927645532</td>
</tr>
<tr>
<td>Document Number:</td>
<td>53-0196965</td>
</tr>
<tr>
<td>Employer Identification Number (EIN):</td>
<td>53-0196965</td>
</tr>
</tbody>
</table>

HHS/ASPR, the primary agency, is responsible for monitoring the work progress of a sub-tasked support agency and approving the request for reimbursement submitted by the support agency. Any costs and supporting documentation must be provided to the HHS/ASPR POC prior to submitting and Intra-governmental Payment and Collection (IPAC). This sub-task will be automatically closed and deobligated 60 days after the project completion date if costs are not presented. As such, all documentation supporting requests for reimbursement shall be submitted to:

David A. Dolinsky, CPA  
Deputy Director for Administration and Finance Operations  
ASPR/Office of Financial Planning and Analysis  
U.S. Department of Health and Human Services  
200 Independence Avenue, SW, Room 638G  
Washington, DC 20201  
202-205-0499 (Office)  
david.dolinsky@hhs.gov
 Accounting Information: (continued)

Reimbursement and billing will be accomplished through the IPAC System.

Program Support Center (PSC), Division of Financial Operations (DFO)
Room 16-30
Parklawn Building
5600 Fishers Lane
Rockville, MD 20857

IPAC will make disbursements/payments.

Document number, ALC, accounting codes, and dollar amounts must be referenced.

<table>
<thead>
<tr>
<th>Authorizing Officials</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>For FDA:</strong></td>
</tr>
<tr>
<td>Signed:</td>
</tr>
<tr>
<td>Name/Title</td>
</tr>
<tr>
<td><strong>For HHS/ASPR:</strong></td>
</tr>
<tr>
<td>Signed:</td>
</tr>
<tr>
<td>David A. Dolinsky</td>
</tr>
<tr>
<td>Deputy Director for Administration and Finance</td>
</tr>
<tr>
<td>Operations/A+F Section Chief</td>
</tr>
<tr>
<td><strong>Funds Certification (ASPR):</strong></td>
</tr>
<tr>
<td>Signed:</td>
</tr>
<tr>
<td>Name/Title</td>
</tr>
</tbody>
</table>
### Appendix D: Characteristics of Potential Foodborne Agents

<table>
<thead>
<tr>
<th>Agent</th>
<th>Incubation Period</th>
<th>Availability</th>
<th>Minimum Infectious Dose, Secondary Transmission</th>
<th>Clinical Syndrome</th>
<th>Case-Fatality</th>
<th>Other Characteristics of Microbe or Illness</th>
</tr>
</thead>
<tbody>
<tr>
<td>Botulinum toxin</td>
<td>Generally between 12 to 72 hours&lt;br&gt;Dependent on the toxin dose, the incubation period varies from 2 hours to 10 days</td>
<td>Organism ubiquitous in environment; cultures need anaerobic conditions</td>
<td>LD₅₀ = 0.001 µg/kg&lt;br&gt;The best estimate of median oral lethal dose for people is 0.2 to 2.0 micrograms</td>
<td>Descending paralysis, respiratory compromise</td>
<td>5%&lt;br&gt;(treated)</td>
<td>95% of patients need hospitalization; 60% of patients need intubation</td>
</tr>
<tr>
<td>Salmonella serotypes (excluding Salmonella typhi)</td>
<td>Can be as short as 6 hours, but is generally 12 to 38 hours</td>
<td>Clinical and research laboratories, culture collections, poultry, environmental sources</td>
<td>10³ organisms; limited secondary transmission</td>
<td>Acute diarrheal illness, 1 to 3% chronic sequelae</td>
<td>&gt;1%</td>
<td>Organism hardy; lengthened survival in the environment</td>
</tr>
<tr>
<td>Salmonella typhi</td>
<td>24 to 72 hours</td>
<td>Clinical and research laboratories</td>
<td>10⁶ organisms; secondary transmission possible&lt;br&gt;Reports exist of illness associated with ingested doses in the range of 100s to 100,000 for S. typhi</td>
<td>Acute febrile illness, protracted recovery, 10% relapse, 1% intestinal rupture</td>
<td>10%&lt;br&gt;untreated, 1% treated</td>
<td>Clinical syndrome unfamiliar in the United States; long incubation period (1 to 3 weeks); produces asymptomatic carrier rate in 3% of cases</td>
</tr>
<tr>
<td>Shigella spp.</td>
<td>24 to 48 hours</td>
<td>Clinical and research laboratories</td>
<td>10² organisms; secondary transmission possible</td>
<td>Acute diarrhea, often bloody</td>
<td>For most common species in the United States, &lt;1%</td>
<td></td>
</tr>
</tbody>
</table>


<table>
<thead>
<tr>
<th>Agent</th>
<th>Incubation Period</th>
<th>Availability</th>
<th>Minimum Infectious Dose, Secondary Transmission</th>
<th>Clinical Syndrome</th>
<th>Case-Fatality</th>
<th>Other Characteristics of Microbe or Illness</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Shigella dysenteriae</em> type 1</td>
<td>12 to 50 hours</td>
<td>Clinical and research laboratories</td>
<td>10 to 100 organisms; secondary transmission possible</td>
<td>Dysentery, seizures</td>
<td>Up to 20%</td>
<td>Causes dysentery, toxic megacolon, hemolytic uremic syndrome (HUS), convulsions in children</td>
</tr>
<tr>
<td><em>Escherichia coli O157:H7</em></td>
<td>24 to 72 hours, but may be as long as 10 days</td>
<td>Clinical and research laboratories, bovine sources, farms</td>
<td>&gt;50 organisms; secondary transmission possible The infectious dose ranges between 1 and 100 cells</td>
<td>Acute bloody diarrhea, 5% HUS, longer term complications</td>
<td>1%</td>
<td>Long-term sequelae: hypertension, stroke, renal insufficiency/ failure neurological complications</td>
</tr>
<tr>
<td><em>Vibrio cholerae</em></td>
<td>24 to 72 hours</td>
<td>Clinical and research laboratories</td>
<td>10⁶ organisms; secondary transmission possible Feeding studies using healthy humans indicated that 1x10⁶ organisms must be ingested to cause illness. Outbreak investigations of natural cholera suggest that, if ingested with water, the infectious dose is 10³ to 10⁶ organisms; if ingested with food, the infectious dose is 10² to 10³ organisms.</td>
<td>Acute life-threatening dehydrating diarrhea</td>
<td>Up to 50% untreated, 1% treated</td>
<td>Historically, causes massive waterborne epidemics in areas with poor sanitation</td>
</tr>
</tbody>
</table>
Appendix E: Chemical Agents – Background Information

Items in this section are as follows:

- General Information
- Classes of Chemical Warfare Agents
- Nerve Agent Exposure Versus Nerve Agent Intoxication
- Chemical Agent Characteristics
- List of Additional Chemical Agents

Additional information on chemical agents may be found at http://emergency.cdc.gov/chemical/.

E.1 GENERAL INFORMATION

Chemical agents are:

- Generally liquid (when containerized).
- Normally disseminated as aerosol or gas.
- Present a respiratory, dermal, ocular, and mucus membrane hazard.
- May be detectable by smell or other senses, but not consistent for all agents.
- Dispersal or lack thereof by weather conditions impacts hazard of agent.

E.2 CLASSES OF CHEMICAL WARFARE AGENTS

**Lethal**

- **Industrial Chemicals**
  - **Choking Agents (Pulmonary Agents).** Choking agents primarily attack lung tissue, causing pulmonary edema. However, there may also be mild irritation of the eyes and upper respiratory tract.
  - **Blood Agents (Cyanides).** Blood agents enter the body following ingestion, skin contact, or inhalation. Once in the body, blood agents inactivate a critical enzyme (cytochrome oxidase); this prevents the cell from using oxygen. Symptoms of blood agent poisoning include respiratory changes, convulsions, and death.

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54 Access additional training in this area via the online courses offered by ORA University at http://intranet.ora.fda.gov/dhrd/default.htm.
• **Warfare Agents**
  - **Blister Agents (Vesicants).** Blister agents include the mustards, arsenicals (primarily Lewisite), and phosgene oxime. They cause pain and tissue damage. Mustards can have delayed effects (hours), but Lewisite and phosgene oxime cause immediate effects. Vesicants cause either fluid-filled blisters or red, elevated skin lesions (phosgene oxime).
  - **Nerve Agents.** Nerve agents cause the greatest concern because of their toxicity, rate of action, and ability to enter the body by multiple routes of entry. Nerve agents are extremely fast acting and classically categorized as colorless and odorless. The symptoms of nerve agents include dimness of vision, runny nose, drooling, difficulty breathing/tightness of chest, nausea, vomiting and diarrhea, muscle jerking or twitching, involuntary urination, defecation, coma, and death.

**Incapacitating and Riot Control**

Incapacitating and riot control agents are not primary terrorist threats due to their relatively short duration of effects and minimal toxicity.

**E.3 NERVE AGENT EXPOSURE VERSUS NERVE AGENT INTOXICATION**

“SLUDGE,” bracketed in the box below, is an acronym often used by medical personnel for identifying the symptoms of organophosphate poisoning.
## E.4 CHEMICAL AGENT CHARACTERISTICS

<table>
<thead>
<tr>
<th>Type of Agent</th>
<th>Effects</th>
<th>Onset</th>
<th>First Aid</th>
<th>Skin Decontamination</th>
<th>Detection</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pulmonary TIC:</strong> CG (PFIB, HC)</td>
<td>Dyspnea, coughing</td>
<td>Hours</td>
<td>None</td>
<td>None usually needed</td>
<td>None</td>
</tr>
<tr>
<td><strong>Cyanide:</strong> AC, CK</td>
<td>Loss of consciousness, convulsions, apnea</td>
<td>Seconds</td>
<td>None (nitrite and thiosulfate)</td>
<td>None usually needed</td>
<td>M256A1</td>
</tr>
<tr>
<td><strong>Vesicants:</strong> H, HD, L</td>
<td>Erythema, blisters; irritation of eyes; cough; dyspnea</td>
<td>Hours (immediate pain after L)</td>
<td>None</td>
<td>M291, M258A1, soap and water, 0.5% bleach</td>
<td>M256A1, M8 and M9 papers, CAM, ACADA, FOX, M90</td>
</tr>
<tr>
<td><strong>Vapor:</strong> miosis, rhinorrhea, dyspnea</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Vapor:</strong> seconds</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Liquid:</strong> sweating, vomiting</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Both:</strong> convulsions, apnea</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Nerve:</strong> GA, GB, GD, GF, VX</td>
<td>Miosis, rhinorrhea, dyspnea</td>
<td>Vapor: seconds</td>
<td>MARK I (1 to 3), Diazepam</td>
<td>M291, M258A1, soap and water, 0.5% bleach</td>
<td>M8A1 alarm, M256A1, M8 and M9 papers, CAM, M22 ACADA</td>
</tr>
<tr>
<td><strong>Vapor:</strong> miosis, rhinorrhea, dyspnea</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Liquid:</strong> sweating, vomiting</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Both:</strong> convulsions, apnea</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Incapacitating:</strong> BZ, Agent 15</td>
<td>Mydriasis, increased body temperature; dry mouth and skin; confusion, visual hallucinations</td>
<td>Minutes to hours</td>
<td>Remove from harming themselves or others</td>
<td>Remove outer clothing; water, or soap and water</td>
<td>None</td>
</tr>
<tr>
<td><strong>Riot Control:</strong> CS, CN</td>
<td>Burning, stinging of eyes, nose, airways, skin</td>
<td>Seconds</td>
<td>None</td>
<td>Water</td>
<td>None</td>
</tr>
</tbody>
</table>


The Medical Management of Chemical Casualties Handbook states, “The [above] table is intended to serve as a reminder of the agents, their effects, first-aid measures, detection, and skin decontamination. It is in no way complete, nor is it intended to be complete.” Consult the appropriate Center SMEs for more detailed information.
## E.5 List of Additional Chemical Agents

Available at [https://emergency.cdc.gov/chemical/factsheets.asp](https://emergency.cdc.gov/chemical/factsheets.asp) with links for agent description.

<table>
<thead>
<tr>
<th>Chemical Agent</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abrin</td>
<td>Methyl Isocyanate</td>
</tr>
<tr>
<td>Adamsite (Diphenylaminechloroarsine or DM)</td>
<td>Mustard Gas (H) (Sulfur Mustard)</td>
</tr>
<tr>
<td>Ammonia</td>
<td>Mustard/Lewisite (HL)</td>
</tr>
<tr>
<td>Arsenic</td>
<td>Mustard/T</td>
</tr>
<tr>
<td>Arsine (SA)</td>
<td>Nicotine</td>
</tr>
<tr>
<td>Barium</td>
<td>Nitrogen Mustard (HN-1, HN-2, HN-3)</td>
</tr>
<tr>
<td>Benzene</td>
<td>Opioids</td>
</tr>
<tr>
<td>Brevetoxin</td>
<td>Osmium Tetroxide</td>
</tr>
<tr>
<td>Bromine (CA)</td>
<td>Paraquat</td>
</tr>
<tr>
<td>Bromobenzylcyanide (CA)</td>
<td>Phosgene (CG)</td>
</tr>
<tr>
<td>BZ (3-Quinuclidinyl Benzilate)</td>
<td>Phosgene Oxime (CX)</td>
</tr>
<tr>
<td>Carbon Monoxide</td>
<td>Phosphine</td>
</tr>
<tr>
<td>Chlorine (CL)</td>
<td>Phosphorus, elemental, white or yellow</td>
</tr>
<tr>
<td>Chloroacetophenone (CN)</td>
<td>Potassium Cyanide (KCN)</td>
</tr>
<tr>
<td>Chlorobenzylidenemalononitrile (CS)</td>
<td>Ricin*</td>
</tr>
<tr>
<td>Chloropicrin (PS)</td>
<td>Sarin (GB)</td>
</tr>
<tr>
<td>Chromium</td>
<td>Saxitoxin</td>
</tr>
<tr>
<td>Colchicine</td>
<td>Selenium</td>
</tr>
<tr>
<td>Cyanide</td>
<td>Sesqui Mustard</td>
</tr>
<tr>
<td>Cyanogen Chloride (CK)</td>
<td>Sodium Azide</td>
</tr>
<tr>
<td>Dibenzoxazepine (CR)</td>
<td>Sodium Cyanide (NaCN)</td>
</tr>
<tr>
<td>Diphosgene (DP)</td>
<td>Sodium Monofluoroacetate (compound 1080)</td>
</tr>
<tr>
<td>Distilled Mustard (HD)</td>
<td>Soman (GD)</td>
</tr>
<tr>
<td>Ethylene glycol</td>
<td>Stibine</td>
</tr>
<tr>
<td>Fentanyl and Other Opioids</td>
<td>Strychnine</td>
</tr>
<tr>
<td>Hydrazine</td>
<td>Sulfur Mustard (H) (Mustard Gas)</td>
</tr>
<tr>
<td>Hydrofluoric Acid (Hydrogen Fluoride)</td>
<td>Sulfur Fluoride</td>
</tr>
<tr>
<td>Hydrogen Chloride</td>
<td>Super Warfarin (Long-Acting Anticoagulant)</td>
</tr>
<tr>
<td>Hydrogen Cyanide (AC)</td>
<td>Tabun (GA)</td>
</tr>
<tr>
<td>Hydrogen Fluoride (Hydrofluoric Acid)</td>
<td>Tetrodotoxin</td>
</tr>
<tr>
<td>Lewisite (L, L-1, L-2, L-3)</td>
<td>Thallium</td>
</tr>
<tr>
<td>LSD (Lysergic Acid Diethylamide)</td>
<td>Trichothecene</td>
</tr>
<tr>
<td>Mercury</td>
<td>VX</td>
</tr>
<tr>
<td>Methyl Bromide</td>
<td>White Phosphorus</td>
</tr>
</tbody>
</table>

*May also be considered a biological agent.*

For more information on chemical emergencies, including specific chemical agents, information for the public, and information for first responders, refer to CDC’s Emergency Preparedness and Response website at [https://emergency.cdc.gov/chemical/index.asp](https://emergency.cdc.gov/chemical/index.asp)
For response information for thousands of hazardous materials, including fire and explosion hazards, health hazards, firefighting techniques, cleanup procedures, protective clothing, and chemical properties, refer to www.cameochemicals.noaa.gov/search/simple.
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Appendix F: Biological Agents – Background Information

Items in this section are as follows:

- General Information
- Classes of Biological Warfare Agents
- Epidemiologic Clues That May Signal a Covert Bioterrorism Attack
- Bioterrorism Agents/Diseases by Category

Additional information on bioterrorism may be found at https://emergency.cdc.gov/bioterrorism/

F.1 GENERAL INFORMATION

Biological agents:

- Produce delayed effects.
- Do not penetrate unbroken skin.
- Do not evaporate.
- Undetectable by senses.
- Difficult to detect in the field.
- Have a range of effects.
- Usually obtained from nature.
- Can now be produced synthetically starting with coding sequence.
- May be engineered for increased virulence.
- Have multiple routes of entry.
- Potentially destroyed by environment.
- Some are contagious.
- Virtually impossible to limit spread or deliberate delivery.
- Environmental detection systems (e.g., BioWatch) for some are now operational.
- Countermeasures are available for some agents.
- Quarantine strategies may be implemented in an effort to control a biological contagion.

F.2 CLASSES OF BIOLOGICAL WARFARE AGENTS

Bacteria. Bacteria are living unicellular organisms and are predominantly self-sufficient (with the exception of several intracellular species that require a host cell to survive). Bacteria require varying conditions of light, moisture, pH, and temperature to survive. Bacteria cause disease in humans by

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55 Access additional training in this area via the online courses offered by ORA University at http://intranet.ora.fda.gov/dhrd/default.htm.
invading host tissues, by producing toxins, or both. The diseases they produce often respond to specific antibiotic therapies.

**Viruses.** Viruses need a “host” cell to live and multiply. However, viruses are technically considered living organisms because they contain genetic material. Although environmental detection units utilizing Polymerase Chain Reaction (PCR) technology can detect viruses, they generally would not be able to assess whether or not the virus is viable.

**Toxins.** Toxins are harmful substances produced by bacteria, plants, and animals. These range from animal venoms (e.g., snake or spider bite), to plant toxins (e.g., ricin), to bacteria toxins (e.g., botulinum toxin). Since they are biochemical moieties (e.g., proteins), they do not replicate and (unlike chemical agents) are nonvolatile and unlikely to cause secondary exposures.

### F.3 Epidemiologic Clues That May Signal a Covert Bioterrorism Attack

Adapted below is a CDC list of epidemiologic clues that may signal a biological incident. In addition, FDA may receive information through its adverse event reporting systems, which indicate a possible biological agent contamination. This list provides the opportunity to recognize and respond to early clues that would be important for minimizing the harmful impacts of a biological incident.

Epidemiologic clues are as follows:

- A preliminary positive signal from BioWatch systems.
- A signal from the BioSense system.
- Intelligence.
- Containers of powders or liquids that are in inappropriate places.
- Large numbers of ill persons with similar disease or syndrome.
- Large numbers of unexplained disease, syndrome, or deaths.
- Unusual illness in a population.
- Higher morbidity and mortality than expected with a common disease or syndrome.
- Failure of a common disease to respond to usual therapy.
- Single case of disease caused by an uncommon agent.
- Multiple unusual or unexplained disease entities coexisting in the same patient without other explanation.
- Disease with an unusual geographic or seasonal distribution.
- Multiple atypical presentations of disease agents.
- Similar genetic type among agents isolated from temporally or spatially distinct sources.
- Unusual, atypical, genetically engineered, or antiquated strain of agent.
- Endemic disease with unexplained increase in incidence.
- Simultaneous clusters of similar illness in noncontiguous areas, domestic or foreign.
- Atypical aerosol, food, or water transmission.
- Ill people presenting near the same time.
- Deaths or illness among animals that precedes or accompanies illness or death in humans.
- No illness in people not exposed to common ventilation systems, but illness among those people in proximity to the systems.
### F.4 Bioterrorism Agents/Diseases by Category

This table was created from information from the CDC Bioterrorism Agents/Diseases by category list (available at [https://emergency.cdc.gov/agent/agentlist.asp](https://emergency.cdc.gov/agent/agentlist.asp)).

<table>
<thead>
<tr>
<th>Biological Diseases/Agents</th>
<th>Category A</th>
<th>Category B</th>
<th>Category C</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Definition:</strong> The U.S. public health system and primary healthcare providers must be prepared to address various biological agents, including pathogens that are rarely seen in the United States. High-priority agents include organisms that pose a risk to national security because they:</td>
<td>- Can be easily disseminated or transmitted from person to person; - Result in high mortality rates and have the potential for major public health impact; - Might cause public panic and social disruption; and - Require special action for public health preparedness.</td>
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**Category A Agents/Diseases**
- Anthrax (*Bacillus anthracis*)
- Botulism (*Clostridium botulinum* toxin)
- Plague (*Yersinia pestis*)
- Smallpox (variola major)
- Tularemia (*Francisella tularensis*)
- Viral hemorrhagic fevers (filoviruses [e.g., Ebola, Marburg] and arenaviruses [e.g., Lassa, Machupo])

**Category B Agents/Diseases**
- Brucellosis (*Brucella* species)
- Epsilon toxin of *Clostridium perfringens*
- Food safety threats (e.g., *Salmonella* species, *Escherichia coli* O157:H7, *Shigella*)
- Glanders (*Burkholderia mallei*)
- Melioidosis (*Burkholderia pseudomallei*)
- Psittacosis (Chlamydia psittaci)
- Q fever (*Coxiella burnetii*)
- Ricin toxin from *Ricinus communis* (castor beans)
- Staphylococcal enterotoxin B
- Typhus fever (*Rickettsia prowazekii*)
- Viral encephalitis (alphaviruses [e.g., Venezuelan equine encephalitis, eastern equine encephalitis, western equine encephalitis])
- Water safety threats (e.g., *Vibrio cholerae*, *Cryptosporidium parvum*)

**Category C Agents**
- Emerging infectious disease threats such as Nipah virus and hantavirus

For more information on specific biological agents/diseases, including agent lists, factsheets, questions and answers, references, treatment, laboratory information, and more, refer to the CDC’s Emergency Preparedness and Response website at [https://emergency.cdc.gov/agent/agentlist.asp](https://emergency.cdc.gov/agent/agentlist.asp).
Appendix G:
Oversight of Select Agents and Toxins by HHS/CDC and USDA/Animal and Plant Health Inspection Service
(7 CFR Part 331, 9 CFR Part 121, and 42 CFR Part 73)

G.1 HHS SELECT AGENTS AND TOXINS

- Abrin
- *Bacillus cereus* Biovar *anthracis*
- Botulinum neurotoxins*
- Botulinum neurotoxin producing species of *Clostridium*
- Conotoxins (Short, paralytic alpha conotoxins containing the following amino acid sequence $X_1CCX_2PACGX_3X_4X_5X_6CX_7$)\(^{56}\)
- *Coxiella burnetii*
- Crimean-Congo haemorrhagic fever virus
- Diacetylscirpenol
- Eastern Equine Encephalitis virus\(^{57}\)
- Ebola virus*
- *Francisella tularensis* *
- Lassa fever virus
- *Lujo* virus
- Marburg virus*
- Monkeypox virus\(^{2}\)
- Reconstructed replication competent forms of the 1918 pandemic influenza virus containing any portion of the coding regions of all eight gene segments (Reconstructed 1918 Influenza virus)
- Ricin
- *Rickettsia prowazekii*
- SARS-associated coronavirus (SARS-CoV)
- Saxitoxin
- South American Haemorrhagic Fever viruses:
  - Chapare
  - Guanarito
  - Junin
  - Machupo
  - Sabia
- Staphylococcal enterotoxins A,B,C,D,E subtypes
- T-2 toxin
- Tetrodotoxin
- Tick-borne encephalitis complex (flavi) viruses:
  - Far Eastern subtype
  - Siberian subtype
  - Kyasanur Forest disease virus
  - Omsk hemorrhagic fever virus
- Variola major virus (smallpox virus)*
- Variola minor virus (Alastrim)*
- Yersinia pestis*

\(^{56}\) C = Cysteine residues are all present as disulfides, with the 1st and 3rd Cysteine, and the 2nd and 4th Cysteine forming specific disulfide bridges; The consensus sequence includes known toxins α-MI and α-GI (shown above) as well as α-GIA, Ac1.1a, α-CnIA, α-CnIB; $X_1 =$ any amino acid(s) or Des-X; $X_2 =$ Asparagine or Histidine; $P =$ Proline; $A =$ Alanine; $G =$ Glycine; $X_3 =$ Arginine or Lysine; $X_4 =$ Asparagine, Histidine, Lysine, Arginine, Tyrosine, Phenylalanine or Tryptophan; $X_5 =$ Tyrosine, Phenylalanine, or Tryptophan; $X_6 =$ Serine, Threonine, Glutamate, Aspartate, Glutamine, or Asparagine; $X_7 =$ Any amino acid(s) or Des X and; “Des X” = “an amino acid does not have to be present at this position.” For example, if a peptide sequence were XCCHPA then the related peptide CCHPA would be designated as Des-X.

\(^{57}\) Select agents that meet any of the following criteria are excluded from the requirements of this part: Any low pathogenic strains of avian influenza virus, South American genotype of eastern equine encephalitis virus, west African clade of Monkeypox viruses, any strain of Newcastle disease virus which does not meet the criteria for virulent Newcastle disease virus, all subspecies Mycoplasma capricolum except subspecies capripneumoniae (contagious caprine pleuropneumonia), all subspecies Mycoplasma mycoides except subspecies mycoides small colony (Mmm SC) (contagious bovine pleuropneumonia), and any subtypes of Venezuelan equine encephalitis virus except for Subtypes IAB or IC, provided that the individual or entity can verify that the agent is within the exclusion category.
G.2 **OVERLAP SELECT AGENTS AND TOXINS**

- *Bacillus anthracis*
- *Bacillus anthracis* Pasteur strain
- *Brucella abortus*
- *Brucella melitensis*
- *Brucella suis*
- *Burkholderia mallei*
- *Burkholderia pseudomallei*
- Hendra virus
- Nipah virus
- Rift Valley fever virus
- Venezuelan Equine Encephalitis virus

G.3 **USDA SELECT AGENTS AND TOXINS**

- African horse sickness virus
- African swine fever virus
- Avian influenza virus
- Classical swine fever virus
- Foot-and-mouth disease virus
- Goat pox virus
- Lumpy skin disease virus
- *Mycoplasma capricolum*
- *Mycoplasma mycoides*
- Newcastle disease virus
- Peste des petits ruminants virus
- Rinderpest virus
- Sheep pox virus
- Swine vesicular disease virus

G.4 **USDA PLANT PROTECTION AND QUARANTINE SELECT AGENTS AND TOXINS**

- *Peronosclerospora philippinensis (Peronosclerospora sacchari)*
- *Phoma glycinicola* (formerly *Pyrenochaeta glycines*)
- *Ralstonia solanacearum*
- *Rathayibacter toxicus*
- *Sclerophthora rayssiae*
- *Synchytrium endobioticum*
- *Xanthomonas oryzae*

05/16/2018
*Denotes Tier 1 Agent

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58 A virulent Newcastle disease virus (avian paramyxovirus serotype 1) has an intracerebral pathogenicity index in day-old chicks (Gallus gallus) of 0.7 or greater or has an amino acid sequence at the fusion (F) protein cleavage site that is consistent with virulent strains of Newcastle disease virus. A failure to detect a cleavage site that is consistent with virulent strains does not confirm the absence of a virulent virus.
Appendix H: Domestic and International Traceback/Traceforward

H.1 Domestic

When there is an FDA-regulated product potentially affected by an emergency, it is necessary to determine all distribution of the affected product. After appropriate consultation with others (State and local officials, Centers, and CDC), OEM/OEO, in coordination with the Office of Regional Operations (ORO), initiates a traceback/traceforward investigation.59 The FDA EOC coordinates the tracing (forward and backward) of the distribution of any potentially contaminated FDA-regulated products and initiates seizures or recalls as appropriate. CDC and local or State agencies may provide assistance on the traceback/traceforward.

H.2 International

If FDA determines international distribution of the product occurred, the Office of International Programs (OIP) notifies foreign governments of the situation as appropriate. OIP is also FDA’s lead office for communicating with various international stakeholders, including the U.S. Department of State (DOS), foreign and U.S. embassies, foreign governments (including counterpart foreign agencies), and international organizations. When FDA learns of suspect or actual incidents involving FDA-regulated products received from or sent to foreign countries, the FDA EOC promptly notifies the Deputy Commissioner for International Programs and the Assistant Director for Communications in OIP, who then coordinate with recall personnel and/or other appropriate staff for the notification and exchange of information with the affected countries.

In addition, OIP coordinates with the FDA EOC to provide timely notification with the appropriate degree of urgency to the following offices as necessary and appropriate:

- FDA Center international offices or liaisons
- Embassies of the country or countries involved
- HHS Office of Global Health Affairs
- Desk officers at DOS
- Other relevant U.S. agency international components
- Appropriate international organizations

Epidemiological traceback data provides the basis for enforcement strategies. At the border, this includes procedures to facilitate movement of similar products that are in compliance with the Federal Food, Drug, and Cosmetic Act of 1938 (FD&C Act) (as mandated in Section 302 of the Public Health Security and Bioterrorism Preparedness and Response Act of 2002 [Bioterrorism Act]) or otherwise not considered a serious threat to public health and safety. Epidemiological traceback data provides intelligence that allows FDA to identify historical shipments through the same or similar streams of commerce of implicated products for appropriate follow-up. Follow-up activities include visiting importers and consignees of previous shipments to identify disposition of imported goods and identifying individuals and firms responsible for the sale, purchase, and receipt of implicated products.

The agency will use the Operational and Administrative System for Input Support or the Mission Activity Reporting and Compliance System and may work with the U.S. Customs and Border Protection (CBP),

59 For more information on how FDA conducts traceback investigations, refer to [www.fda.gov/ICECI/Inspections/InspectionGuides/ucm075005.htm](http://www.fda.gov/ICECI/Inspections/InspectionGuides/ucm075005.htm) and FDA’s “Guide to Traceback of Fresh Fruits and Vegetables Implicated in Epidemiological Investigations” (June 2006).
located at the Commercial Trade Analytical Center, to analyze entry data in order to support decisions for dispatching human resources through assignments or other methods established for rapid response to support operations. The Director of the Division of Import Operations and Policy (DIOP) will advise CBP of the situation and the information needed to use CBP’s Automated Targeting System (ATS) to detect and detain future shipments of potentially tainted products before their introduction into U.S. commerce.
Appendix I:
Memorandum of Understanding

A memorandum of understanding (MOU) is a critical component of any formal arrangement of cooperation between two or more entities. An MOU usually describes, in broad general terms, an area of mutual interest or concern that two or more agencies or organizations may cooperatively address. MOUs generally do not include specific information regarding detailed scope of work or exchange of funds or human resources. Below is a table of the current, finalized MOU that FDA has with other organizations.

A complete listing of **MOUs or other written cooperative arrangements** with States, other Federal agencies and foreign government counterparts, and the World Health Organization (WHO) can be found at [www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/default.htm](http://www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/default.htm).

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<th>Understanding or Agreement</th>
<th>Purpose</th>
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<th>Notes/Dates (Effective/Expiration)</th>
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<td>FDA and CDC</td>
<td><a href="https://www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/DomesticMOUs/ucm402130.htm">https://www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/DomesticMOUs/ucm402130.htm</a></td>
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<td><a href="http://www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/DomesticMOUs/ucm294512.htm">www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/DomesticMOUs/ucm294512.htm</a></td>
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<td>225-72-2009</td>
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<td>FDA and USDA Agricultural Marketing Service (AMS)</td>
<td><a href="http://www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/DomesticMOUs/ucm115864.htm">www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/DomesticMOUs/ucm115864.htm</a></td>
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<td>225-12-007</td>
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<td>FDA and USDA</td>
<td><a href="https://www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/DomesticMOUs/ucm294512.htm">https://www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/DomesticMOUs/ucm294512.htm</a></td>
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<td>Exchange of information</td>
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<td><a href="https://www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/DomesticMOUs/ucm294512.htm">https://www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/DomesticMOUs/ucm294512.htm</a></td>
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<td>N/A</td>
<td>Sharing and exchange of Information</td>
<td>FDA and Health Products and Food Branch (HPFB), Health Canada of Canada</td>
<td><a href="http://www.fda.gov/InternationalPrograms/Agreements/MemorandaofUnderstanding/ucm197638.htm">www.fda.gov/InternationalPrograms/Agreements/MemorandaofUnderstanding/ucm197638.htm</a></td>
<td>04/16/2008 11/18/2013</td>
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<tr>
<td>225-96-4001</td>
<td>Trilateral: cooperation in scientific and regulatory fields of health</td>
<td>FDA; Health Protection Branch (HPB), Health Canada of Canada; and Subsecretaría de Regulación y Fomento Sanitario of the Secretaría de Salud (SSA) of the United Mexican States</td>
<td><a href="http://www.fda.gov/InternationalPrograms/Agreements/MemorandaofUnderstanding/ucm107612.htm">www.fda.gov/InternationalPrograms/Agreements/MemorandaofUnderstanding/ucm107612.htm</a></td>
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*USDA Farm and Foreign Agricultural Services (FFAS)/Food, Nutrition, and Consumer Services (FNCS)/Food Safety (FS)/Marketing and Regulatory Programs (MRP)/Research, Education, and Economics (REE)
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Listed below are some common forms and templates to be used by FDA personnel during the course of an incident. Hyperlinks attached to each title link directly to the appropriate form/template included within this appendix. Electronic copies can be found on the EON-IMS.

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<thead>
<tr>
<th>Title</th>
<th>Page #</th>
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<tbody>
<tr>
<td>Incident Briefing (ICS 201), Adapted for FDA</td>
<td>App J-3</td>
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<tr>
<td>Incident Objectives (ICS 202), Adapted for FDA</td>
<td>App J-7</td>
</tr>
<tr>
<td>Organization Assignment List (ICS 203), Adapted for FDA</td>
<td>App J-9</td>
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<tr>
<td>Assignment List (ICS 204), Adapted for FDA</td>
<td>App J-11</td>
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<tr>
<td>Incident Communications Plan (ICS 205), Adapted for FDA</td>
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<td>Medical Plan (ICS 206), Adapted for FDA</td>
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<td>Safety Message/Plan (ICS 208)</td>
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<td>Incident Check-In List (ICS 211), Adapted for FDA</td>
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<td>Resource Request (ICS 213 RR), Adapted for FDA</td>
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<tr>
<td>Activity Log (ICS 214)</td>
<td>App J-23</td>
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<td>Operational Planning Worksheet (ICS 215)</td>
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<td>Demobilization Check-Out (ICS 221), Adapted for FDA</td>
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<tr>
<td>Situation Report</td>
<td>App J-31</td>
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## Incident Briefing (ICS 201), Adapated for FDA

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<th>Incident Name:</th>
<th>Incident Number: (EON num. if applicable)</th>
<th>Date/Time Initiated: Date: Time:</th>
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4. **Map/Sketch** (include sketch, showing the total area of operations, the incident site/area, impacted and threatened areas, overflight results, trajectories, impacted shorelines, or other graphics depicting situational status and resource assignment):

5. **Situation Summary and Health and Safety Briefing** (for briefings or transfer of command): Recognize potential incident Health and Safety Hazards and develop necessary measures (remove hazard, provide personal protective equipment, warn people of the hazard) to protect responders from those hazards.

6. **Prepared by**: Name: Position/Title: Signature:  
ICS 201, Page 1  
Date/Time: 

Updated by FDA 2/2011
<table>
<thead>
<tr>
<th>1. Incident Name:</th>
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7. Current and Planned Objectives:

8. Current and Planned Actions, Strategies, and Tactics:

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3. Date/Time Initiated: Date: Time: |

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Updated by FDA 2/2011

1. Incident Name: |
2. Incident Number: (EON num. if applicable) |
3. Date/Time Initiated: Date: Time: |

9. Current Organization (fill in additional organization as appropriate):
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ICS 201, Page 4

Updated by FDA 2/2011
# Incident Objectives (ICS 202), Adapted for FDA

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<td>☐ Map/Chart</td>
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<td>☐ ICS 204</td>
<td>Weather Forecast/Tides/Currents</td>
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<th>Name: ___________ Position/Title: ___________ Signature: ___________</th>
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</thead>
</table>

<table>
<thead>
<tr>
<th>Approved by Incident Commander:</th>
<th>Name: ___________ Signature: ___________</th>
</tr>
</thead>
</table>

ICS 202  IAP Page _____  Date/Time: ___________

Updated by FDA 2/2011
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# Organization Assignment List (ICS 203), Adapted for FDA

1. **Incident Name:**

2. **Operational Period:**
   - Date From: 
   - Date To: 
   - Time From: 
   - Time To: 

3. **Incident Commander(s)/Agency Incident Coordinator and Command Staff:**
   - IMT IC/UCs: Chief
   - IMG AIC: Deputy

4. **Agency/Organization Representatives:**

5. **Planning Section:**
   - Chief
   - Deputy
   - Resource Unit
   - Situation Unit
   - Documentation Unit
   - Demobilization Unit
   - Technical Specialists

6. **Logistics Section:**
   - Chief
   - Deputy
   - Support Branch
   - Director
   - Supply Unit
   - Facilities Unit

7. **Operations Section:**

8. **Finance/Administration Section:**
   - Ground Support Unit: Chief
   - Service Branch: Deputy
   - Director
   - Communications Unit
   - Medical Unit
   - Food Unit

9. **Prepared by:**
   - Name: 
   - Position/Title: 
   - Signature: 

ICS 203

Date/Time: 

Updated by FDA 2/2011

---

**Appendix J:**

July 2019

App J-10

Forms and Templates
This Page Is Intentionally Left Blank.
# Assignment List (ICS 204), Adapated for FDA

## 1. Incident Name: ___

## 2. Operational Period:
- **Date From:** ___
- **Date To:** ___
- **Time From:** ___
- **Time To:** ___

## 3. Branch: ___
**Division:** ___
**Group:** ___

## 4. Operations Personnel:

<table>
<thead>
<tr>
<th>Name</th>
<th>Contact Number(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operations Section Chief</td>
<td></td>
</tr>
<tr>
<td>Branch Director</td>
<td></td>
</tr>
<tr>
<td>Division/Group Supervisor</td>
<td></td>
</tr>
</tbody>
</table>

## 5. Resources Assigned:

<table>
<thead>
<tr>
<th>Resource Identifier</th>
<th>Leader</th>
<th># of Persons</th>
<th>Contact (e.g., phone, pager, radio frequency, etc.)</th>
<th>Reporting Location, Special Equipment and Supplies, Remarks, Notes, Information</th>
</tr>
</thead>
<tbody>
<tr>
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</tbody>
</table>

## 6. Work Assignments:


## 7. Special Instructions:


## 8. Communications (radio and/or phone contact numbers needed for this assignment):

<table>
<thead>
<tr>
<th>Name/Function</th>
<th>Primary Contact: indicate cell, pager, or radio (frequency/system/channel)</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

## 9. Prepared by:
- **Name:** ___
- **Position/Title:** ___
- **Signature:** ___

---

Updated by FDA 2/2011
This Page Is Intentionally Left Blank.
# Incident Communications Plan (ICS 205), Adapted for FDA

## 1. Incident Name:

## 2. Date/Time Prepared:

<table>
<thead>
<tr>
<th>Date:</th>
<th>Time:</th>
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</table>

## 3. Operational Period:

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<tr>
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<th>Date To:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time From:</td>
<td>Time To:</td>
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</tbody>
</table>

## 4. Incident communication information:

<table>
<thead>
<tr>
<th>Incident Assigned Position</th>
<th>Name (Last, First)</th>
<th>Primary Number</th>
<th>Secondary Number</th>
<th>Other Method (s) of Contact (pager, email, radio, etc.)</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

## 5. Special Instructions:

## 6. Prepared by (Communications Unit Leader):

<table>
<thead>
<tr>
<th>Name:</th>
<th>Signature:</th>
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<tbody>
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**ICS 205**

**IAP Page _____**

**Date/Time:**

---

*Updated by FDA 2/2011*
This Page Is Intentionally Left Blank.
## Medical Plan (ICS 206), Adapted for FDA

### 1. Incident Name: [ ]

### 2. Operational Period:

<table>
<thead>
<tr>
<th>Date From</th>
<th>Date To</th>
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<tbody>
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<table>
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<tr>
<th>Time From</th>
<th>Time To</th>
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<tbody>
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### 3. Medical Aid Stations:

<table>
<thead>
<tr>
<th>Name</th>
<th>Location</th>
<th>Contact Number(s)/Frequency</th>
<th>Paramedics on Site?</th>
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<tbody>
<tr>
<td></td>
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<td>Yes</td>
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<td>Yes</td>
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</table>

### 4. Transportation:

<table>
<thead>
<tr>
<th>Ambulance Service</th>
<th>Location</th>
<th>Contact Number(s)/Frequency</th>
<th>Level of Service</th>
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<tbody>
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<td>ALS</td>
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### 5. Hospitals:

<table>
<thead>
<tr>
<th>Hospital Name</th>
<th>Address</th>
<th>Contact Number(s)</th>
<th>Distance</th>
<th>Trauma Center</th>
<th>Burn Center</th>
<th>Helipad</th>
</tr>
</thead>
<tbody>
<tr>
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<td>Yes</td>
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<td>Yes</td>
<td>No</td>
<td>Yes</td>
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</tbody>
</table>

### 6. Special Medical Emergency Procedures:

<p>| | | | | | | | | |</p>
<table>
<thead>
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</thead>
</table>

### 7. Prepared by (Medical Unit Leader):

Name: ___________________________ Signature: ___________________________

### 8. Approved by (Safety Officer):

Name: ___________________________ Signature: ___________________________

ICS 206       IAP Page ______   Date/Time: ___________________________

Updated by FDA 2/2011

---

July 2019

Appendix J: Forms and Templates
### Safety Message/Plan (ICS 208)

<table>
<thead>
<tr>
<th>1. Incident Name:</th>
<th>2. Operational Period: Date From:</th>
<th>Date To:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Time From:</td>
<td>Time To:</td>
</tr>
</tbody>
</table>


4. Site Safety Plan Required? Yes ☐ No ☐
   Approved Site Safety Plan(s) Located At:

5. Prepared by: Name: _______________ Position/Title: _______________ Signature: _______________

ICS 208 | IAP Page ____ | Date/Time: _______________
### Incident Check-In List (ICS 211), Adapted for FDA

<table>
<thead>
<tr>
<th>1. Organization Type</th>
<th>Check one:</th>
<th>2. Incident Name</th>
<th>3. Check-in Location</th>
<th>4. Date/Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>IMT</td>
<td></td>
<td></td>
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<tr>
<td>IMG</td>
<td></td>
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<tr>
<td>________________</td>
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#### Check-In Information

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</tbody>
</table>

15. Prepared by (Name and Position). Use back for additional remarks or comments.

Page ___ of ___

Updated by FDA 2/2011

July 2019

App J-20

Appendix J:
Forms and Templates
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# Resource Request (ICS 213 RR), Adapted for FDA

<table>
<thead>
<tr>
<th>1. Incident Name:</th>
<th>2. Date/Time</th>
<th>3. Resource Request Number:</th>
</tr>
</thead>
</table>

### 4. Order
(Use additional forms when requesting different resource sources of supply.)

<table>
<thead>
<tr>
<th>Qty.</th>
<th>Kind</th>
<th>Type</th>
<th>Detailed Item Description: (Vital characteristics, brand, specs, experience, size, etc.)</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
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</tbody>
</table>

### 5. Resource Status

<table>
<thead>
<tr>
<th>Received by</th>
<th>Date/Time</th>
<th>Assigned to</th>
<th>Released to</th>
<th>Date/Time</th>
</tr>
</thead>
<tbody>
<tr>
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</tbody>
</table>

### 6. Requested Delivery/Reporting Location:

### 7. Suitable Substitutes and/or Suggested Sources:

### 8. Requested by Name/Position: 9. Priority: ☐ Urgent  ☐ Routine  ☐ Low

### 10. Section Chief Approval:

### 11. Logistics Order Number:

### 12. Supplier Phone/Fax/Email:

### 13. Name of Supplier/POC:

### 14. Notes:

### 15. Approval Signature of Auth Logistics Rep: 16. Date/Time:

### 17. Order placed by:

### 18. Reply/Comments from Finance:

### 19. Finance Section Signature: 20. Date/Time:

---

ICS 213 RR, Page 1

Updated by FDA 2/2011

---
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## Activity Log (ICS 214)

<table>
<thead>
<tr>
<th>1. Incident Name:</th>
<th>2. Operational Period: Date From:</th>
<th>Date To:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Time From:</td>
<td>Time To:</td>
</tr>
</tbody>
</table>

|----------|------------------|---------------------------|

<table>
<thead>
<tr>
<th>6. Resources Assigned:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
</tr>
<tr>
<td>------</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>7. Activity Log:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date/Time</td>
</tr>
<tr>
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</table>

8. Prepared by: Name: __________________________ Position/Title: __________________________ Signature: __________________________

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ICS 214, Page 1
## Operational Planning Worksheet (ICS 215)

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<tbody>
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<td></td>
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<td>Req.</td>
<td>Have</td>
<td>Need</td>
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<td>Name: ______________________</td>
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<td>Req.</td>
<td>Have</td>
<td>Need</td>
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<td>Position/Title: ______________</td>
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<td>Have</td>
<td>Need</td>
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<td>Signature: __________________</td>
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<td>Req.</td>
<td>Have</td>
<td>Need</td>
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<td>Date/Time: __________________</td>
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ICS 215
Incident Action Plan Safety Analysis (ICS 215A)

<table>
<thead>
<tr>
<th>1. Incident Name:</th>
<th>2. Incident Number:</th>
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</thead>
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<table>
<thead>
<tr>
<th>3. Date/Time Prepared:</th>
<th>4. Operational Period: Date From: Date To:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date:</td>
<td>Time: Time From: Time To:</td>
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</table>

8. Prepared by (Safety Officer): Name: __________________________ Signature: __________________________

Prepared by (Operations Section Chief): Name: __________________________ Signature: __________________________

ICS 215A Date/Time: __________________________
This Page Is Intentionally Left Blank.
## Demobilization Check-Out (ICS 221), Adapted for FDA

<table>
<thead>
<tr>
<th>1. Incident Name:</th>
<th>2. Incident (EON) Number:</th>
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</thead>
<tbody>
<tr>
<td></td>
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</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>3. Planned Release Date/Time:</th>
<th>4. Resource or Personnel Released:</th>
<th>5. Order Request Number:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date</td>
<td>Time</td>
<td></td>
</tr>
</tbody>
</table>

### 6. Resource or Personnel: [ ] IMT  [ ] IMG

You and your resources are in the process of being released. Resources are not released until the checked boxes below have been signed off by the appropriate overhead and the Demobilization Unit Leader (or Planning Section representative).

#### OPERATIONS SECTION

<table>
<thead>
<tr>
<th>Division/Group Supervisor</th>
<th>Remarks</th>
<th>Name</th>
<th>Signature</th>
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#### PLANNING SECTION

<table>
<thead>
<tr>
<th>Unit/Leader</th>
<th>Remarks</th>
<th>Name</th>
<th>Signature</th>
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#### LOGISTICS SECTION

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#### FINANCE/ADMINISTRATION SECTION

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#### OTHER SECTION/STAFF

<table>
<thead>
<tr>
<th>Unit/Other</th>
<th>Remarks</th>
<th>Name</th>
<th>Signature</th>
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### 7. Remarks:

### 8. Travel Information:

<table>
<thead>
<tr>
<th>Estimated Time of Departure:</th>
<th>Actual Release Date/Time:</th>
<th>Room Overnight: [ ] Yes  [ ] No</th>
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<tbody>
<tr>
<td>Destination</td>
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<td>Travel Method:</td>
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<td>Departure Location:</td>
<td>Office/Center/Region/District Notified:</td>
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<tr>
<td>Reservation, Flight, or Train Number:</td>
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### 9. Reassignment Information: [ ] Yes  [ ] No

<table>
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<th>Incident Name:</th>
<th>Incident Number:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Location:</td>
<td>Order Request Number:</td>
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### 10. Prepared by:

<table>
<thead>
<tr>
<th>Name:</th>
<th>Position/Title:</th>
<th>Signature:</th>
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ICS 221 Date/Time: ____________________________

Updated by FDA 9/2013
## FDA Situation Report

### Page Header

**Distributing Office (EOC)**

<table>
<thead>
<tr>
<th>Title of Incident</th>
<th>FDA Situation Report Number</th>
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<tr>
<th>Date (DD/MM/YYYY)</th>
<th>Timeframe Report Covers (military time MM/DD/YYYY to MM/DD/YYYY)</th>
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### A. Executive Summary:

- A brief description of the incident
- Initial and long-term response operations
- The resources that were deployed, including whether specific resources were requested through the SOC, FEMA, mutual aid, or another mechanism
- A description of key events and the timeframes of occurrence
- Decisions that were made in response to events and the results
- Issues that arose during the course of operations. Include those between or among MAC System entities, between the Incident Command and the EOC, and within the EOC.

### B. New Information:

### C. Investigations-Related Information and Other Data:

Data sources may include local, State, and other Federal Departments/Agencies. The following are examples of what data to report in this section.

- Outbreak Data
- Epidemiological Data
- Lab Testing Data
- Analytical Data
  - Traceback Data
  - Traceforward Data
  - Assignments
- Sampling Data

### D. Documentation Unit:
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E. Communications:
- Internal (within FDA)
- External (with other agencies, including HHS and CDC)
- Public Information (issuance of press releases; Consumer Health Information posted to FDA website; Call Center; outreach to patient, health professional and industry groups)

F. Strategic Objectives:

G. Additional Incident Response Information:

H. Prepared by:

I. Reviewed/Approved by:
## Appendix K:
### Glossary of Terms

This appendix contains definitions of key terms as they are applied within the FDA EOP. All supporting agency emergency plans and procedural documentation shall be compliant with the terminology contained herein.

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
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<tbody>
<tr>
<td>Abbreviated New Drug Application (ANDA)</td>
<td>An ANDA contains data that, when submitted to FDA’s Center for Drug Evaluation and Research, Office of Generic Drugs, provide for the review and ultimate approval of a generic drug product. Once approved, an applicant may manufacture and market the generic drug product to provide a safe, effective, low-cost alternative to the American public. A generic drug product is one that is comparable to an innovator drug product in dosage form, strength, route of administration, quality, performance characteristics, and intended use. Generic drug applications are termed “abbreviated” because they are generally not required to include preclinical (animal) and clinical (human) data to establish safety and effectiveness. Instead, generic applicants must scientifically demonstrate that their product is bioequivalent (i.e., performs in the same manner as the innovator drug). One way scientists demonstrate bioequivalence is to measure the time it takes the generic drug to reach the bloodstream in 24 to 36 healthy volunteers. This gives them the rate of absorption, or bioavailability, of the generic drug, which they can then compare to that of the innovator drug. The generic version must deliver the same amount of active ingredients into a patient’s bloodstream in the same amount of time as the innovator drug. The ANDA regulations can be found in Title 21 of the CFR Part 314, Subpart C.</td>
</tr>
<tr>
<td>Activation</td>
<td>Implementation of an emergency plan, in whole or in part, in response to a confirmed report of an incident.</td>
</tr>
<tr>
<td>Agency Emergency Coordinator</td>
<td>Employees who serve as FDA’s central emergency coordination point with FDA field offices, headquarters, and Centers and Federal, State, and local entities, including the CDC. They provide interagency coordination and response during adverse events, foodborne illnesses, injuries, product tampering, and manmade and natural disasters. In addition, they provide essential support and lead the agency proportional response to bioterrorism, shortage, and recall emergencies. They serve as a SME in the Incident Command System (ICS) and the National Incident Management System (NIMS).</td>
</tr>
<tr>
<td>Agency Executive Group (AEG)</td>
<td>The AEG is established when an emergency calls for the involvement of senior FDA officials with the knowledge and authority to address a wide range of issues that may arise. The AEG, whether convened as a collective unit or acting as individual members, serves primarily to provide strategic policy direction and guidance for major agency emergency response activities and to approve important policy decisions in consultation with the Commissioner and AIC when necessary.</td>
</tr>
<tr>
<td>Agency Incident Coordinator (AIC)</td>
<td>The individual responsible for the overall coordination of agency emergency management teams assigned. The AIC ensures that conflicts are resolved, incident objectives are established, and strategies are selected for the use of critical agency resources. He/she is also responsible for communicating with FDA headquarters and field organizational components and external Federal agencies. The AIC reports directly to the AEG. Until an AIC is appointed by the Commissioner, Deputy Commissioner, or other designated senior agency official, the Director, OEO, within OEM, may serve as the AIC.</td>
</tr>
</tbody>
</table>
| Alert | Incident information without support. An alert is issued to agency emergency personnel when the following is received:  
- An unconfirmed report of a product-related illness/injury or unanticipated adverse reaction.  
- An unconfirmed report of the presence of a toxic (i.e., chemical, biological, radiological, or nuclear [CBRN]) substance.  
- A report of a natural (e.g., hurricane, flood, tornado) or manmade (e.g., oil spill, radiological accident) disaster. |
| All-Hazards | Describing an incident, natural or manmade, that warrants action to protect life, property, environment, and public health or safety and to minimize disruptions of government, social, or economic activities. |
| Antiviral | Drug that is used to prevent or cure a disease caused by a virus by interfering with the ability of the virus to multiply in number or spread from cell to cell. |
| **Avian Influenza** | An infection caused by type A influenza viruses. These viruses occur naturally among birds worldwide, particularly waterfowl and shore birds. Wild birds are the natural reservoir of avian influenza viruses and carry the viruses in their intestines; they usually do not become ill from them. Infected birds shed influenza virus in their saliva, nasal secretions, and feces. Susceptible birds become infected when they have contact with contaminated excretions or with surfaces that are contaminated with excretions or secretions. Domesticated poultry may become infected with avian influenza virus through direct contact with infected waterfowl or other infected poultry or through contact with surfaces (such as dirt or cages) or materials (such as water or feed) that have been contaminated with the virus. |
| **Becquerel (Bq)** | A unit of radioactivity equivalent to one decay (disintegration) per second. |
| **Catastrophic Incident** | Any natural or manmade incident, including terrorism, that results in extraordinary levels of mass casualties, damage, or disruption severely affecting the population, infrastructure, environment, economy, national morale, and/or government functions. |
| **Center/Office Emergency Coordinator** | The staff member who has responsibility for Center/Office emergency management programs and activities. The role is one of coordinating all aspects of the Center’s mitigation, preparedness, response, and recovery capabilities and serving in a liaison capacity between the Center/Office emergency management team and the Operations Section Chief. The Center Emergency Coordinator may serve as the AIC for all emergencies and disasters involving or affecting his/her product Center, unless otherwise directed. |
| **Centers** | FDA Product Centers:  
- Center for Biologics Evaluation and Research (CBER)  
- Center for Drug Evaluation and Research (CDER)  
- Center for Devices and Radiological Health (CDRH)  
- Center for Food Safety and Applied Nutrition (CFSAN)  
- Center for Veterinary Medicine (CVM)  
- Center for Tobacco Products (CTP)  
FDA Research Center:  
- National Center for Toxicological Research (NCTR) |
<p>| <strong>Centers for Disease Control and Prevention (CDC)</strong> | The U.S. government agency at the forefront of public health efforts to prevent and control infectious and chronic diseases, injuries, workplace hazards, disabilities, and environmental health threats. CDC is one of 13 major operating components of HHS. |
| <strong>Chief</strong> | The ICS title for individuals responsible for management of functional Sections: Operations, Planning, Logistics, and Finance/Administration. |
| <strong>Command Staff</strong> | The staff who report directly to the ICAIC, including the Public Information Officer (PIO), Safety Officer, Liaison Officer (LNO), and other positions as required. They may have an assistant or assistants as needed. |
| <strong>Commissioner</strong> | The Commissioner of Food and Drugs. |
| <strong>Committed Dose</strong> | The cumulative dose to an individual’s whole body or a specific organ resulting from continuous exposure to radioactive materials deposited inside the body, projected to either 50 or 70 years for an occupational worker or member of the general public, respectively. |
| <strong>Common Operating Picture</strong> | A continuously updated overview of an incident compiled throughout an incident’s lifecycle from data shared between integrated systems for communication, information management, and intelligence and information sharing. The common operating picture allows agency headquarters and field personnel to make effective, consistent, and timely decisions. |
| <strong>Confirmed Report</strong> | Report that a problem has been confirmed through laboratory analyses, field investigations, analysis of epidemiological data, or a combination of these. Information received from another governmental agency or other source known to be reliable may be accepted for confirmation purposes. |
| <strong>Contamination (Radioactive)</strong> | The unexpected or undesired presence of radioactive material in or on the surfaces of structures, areas, objects, or people. |
| <strong>Cooperating Agency</strong> | Cooperating agencies include Federal agencies that provide additional technical and resource support specific to nuclear/radiological incidents to DHS and the coordinating agencies. |
| <strong>Coordinating Agency</strong> | Coordinating agencies provide the leadership, expertise, and authorities to implement critical and specific nuclear/radiological aspects of the response, and facilitate nuclear/radiological aspects of the response in accordance with those authorities and capabilities. The coordinating agencies are those Federal agencies that own, have custody of, authorize, regulate, or are otherwise assigned responsibility for the nuclear/radioactive material, facility, or activity involved in the incident. These Federal agencies have nuclear/radiological authorities, technical expertise, and/or assets for responding to the unique characteristics of nuclear/radiological incidents that are not otherwise described in the NRF. |
| <strong>Corrective Action</strong> | Implementing procedures that are based on lessons learned from actual incidents or from training and exercises. |
| <strong>Decay (Radioactive)</strong> | Disintegration of the nucleus of an unstable atom by the release of radiation. |
| <strong>Demobilization</strong> | The orderly, safe, and efficient return of agency resources to their original location and operating status. Demobilization signals the transition from the response phase of operations to recovery. |
| <strong>Deployment</strong> | The relocation of agency resources to desired operational areas. Deployment encompasses all activities from departing normal locations through arrival at destinations. |
| <strong>Division</strong> | The organizational level having responsibility for operations within a defined geographic area. Divisions are established within the Operations Section, at the headquarters level, to coordinate and monitor the operations of responding FDA Program and/or Districts within a particular location and/or when a foreign country is involved. |
| <strong>Dose Coefficient</strong> | A factor used to convert radionuclide intake or content to radiation dose. One usually expresses it as dose per unit intake (e.g., sieverts [Svs] per becquerel). |
| <strong>Dose Equivalent</strong> | A quantity used in radiation protection to place all radiation on a common scale for calculating tissue damage. Dose equivalent is the absorbed dose in grays times a quality factor. The quality factor accounts for differences in radiation effects caused by different types of ionizing radiation. Some radiation, including alpha particles, causes a greater amount of damage per unit of absorbed dose than other radiation. The Sv is the unit that measures dose equivalent. |
| <strong>Dosimeter</strong> | A small portable instrument (e.g., a film badge, thermoluminescent dosimeter [TLD], or pocket dosimeter) for measuring and recording the total accumulated dose of ionizing radiation a person receives. |
| <strong>Effective Dose</strong> | A dosimetric quantity useful for comparing the overall health effects of irradiation of the whole body. It takes into account the absorbed doses received by various organs and tissues and weights them according to present knowledge of the sensitivity of each organ to radiation. It also accounts for the type of radiation and the potential for each type to inflict biologic damage. Use the effective dose, for example, to compare the overall health detrimental of different radionuclides in a given mix. The unit of effective dose is the Sv; 1 Sv = 1 joule per kilogram (J/kg). |
| <strong>Emergency</strong> | Any incident, whether natural or manmade, that requires responsive action to protect life or property. Under the Stafford Act, an emergency means any occasion or instance for which, in the determination of the President, Federal assistance is needed to supplement State and local efforts and capabilities to save lives and to protect property and public health and safety, or to lessen or avert the threat of a catastrophe in any part of the United States. |
| <strong>Emergency Management</strong> | The coordination and integration of all activities necessary to build, sustain, and improve the capability to prepare for, protect against, respond to, recover from, or mitigate against threatened or actual natural disasters, acts of terrorism, or other manmade emergencies. |
| <strong>Emergency Operations Plan (EOP)</strong> | An ongoing plan for responding to a wide variety of potential hazards. |
| <strong>Emergency Support Function (ESF)</strong> | Used by the Federal government and many State governments as the primary mechanism at the operational level to organize and provide assistance. ESFs align categories of resources and provide strategic objectives for their use. ESFs utilize standardized resource management concepts such as typing, inventoring, and tracking to facilitate the dispatch, deployment, and recovery of resources before, during, and after an incident. |
| <strong>Emergency Support Function (ESF) Annexes</strong> | Annexes to the FDA EOP that present the missions, policies, structures, and responsibilities of FDA organizational components for coordinating resource and programmatic support to other Federal agencies, States, territories, tribes, and local jurisdictions when NRF ESFs are activated during an incident. |</p>
<table>
<thead>
<tr>
<th>Term</th>
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<tr>
<td>Emergency Use Authorization (EUA)</td>
<td>Section 564 of the FD&amp;C Act (21 U.S.C. 360bbb-3) permits the Commissioner of Food and Drugs to authorize the use of an unapproved medical product or an unapproved use of an approved medical product during a declared emergency involving a heightened risk of attack on the public or U.S. military forces. The EUA authority allows FDA to strengthen the public health protections against CBRN agents that may be used to attack the American people or U.S. Armed Forces. Under Section 564 of the FD&amp;C Act, the FDA Commissioner may allow medical countermeasures to be used in an emergency to diagnose, treat, or prevent serious or life-threatening diseases or conditions caused by such agents when there are no adequate, approved, and available alternatives. Before FDA can issue an EUA, the HHS Secretary must declare an emergency justifying the authorization to use the product, based on one of three determinations: (1) a determination by the Secretary of Homeland Security of a domestic emergency, or the significant potential for a domestic emergency; (2) a determination by the Secretary of Defense of a military emergency, or the significant potential for a military emergency; or (3) a determination by the HHS Secretary of a public health emergency under the Public Health Service Act (PHSA).</td>
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<tr>
<td>Epidemic</td>
<td>A disease occurring suddenly in humans in a community, region, or country in numbers clearly in excess of normal. (See Pandemic.)</td>
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<tr>
<td>Exposure (Radiation)</td>
<td>A measure of ionization in air caused by x-rays or gamma rays only. The unit of exposure most often used is the roentgen (R).</td>
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<tr>
<td>Exposure Pathway</td>
<td>A route by which a radionuclide or other toxic material can enter the body. The main exposure routes are inhalation, ingestion, absorption through the skin, and entry through a cut or wound in the skin.</td>
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<tr>
<td>Finance/Administration Section</td>
<td>The ICS Section responsible for developing cost analyses and ensuring the IAP is within financial limits established by the Incident Commander. This Section also develops contracts and pays for resources.</td>
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<tr>
<td>Good Manufacturing Practice (GMP)</td>
<td>Manufacturers establish and follow quality systems to help ensure their products consistently meet applicable requirements and specifications. The quality systems for FDA-regulated products (food, drugs, biological products, and devices) are known as current GMPs, also seen as cGMP or CGMP. The GMP regulations for food are at 21 CFR Part 110. For human drugs and most biological products, the CGMP regulations are at 21 CFR Parts 210 and 211. For devices, the CGMP regulations are at 21 CFR Part 820 (quality system regulation).</td>
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<tr>
<td>Group</td>
<td>An organizational subdivision established to divide the incident management structure into functional areas of operation. Groups within the Operations Section, composed of headquarters personnel, are created to address similar emergency response activities.</td>
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<td>Hazard</td>
<td>Something that is potentially dangerous or harmful, often the root cause of an unwanted outcome.</td>
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<tr>
<td>Headquarters</td>
<td>Includes all Offices within the OC, ORA, product Centers, and any divisions or branches therein.</td>
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<tr>
<td>Improvised Nuclear Device (IND)</td>
<td>An IND is an illicit nuclear weapon bought, stolen, or otherwise originating from a nuclear state, or a weapon fabricated by a terrorist group from illegally obtained fissile nuclear weapons material that produces a nuclear explosion. The nuclear yield achieved by an IND produces extreme heat, powerful shockwaves, and prompt radiation that would be acutely lethal for a significant distance. It also produces radioactive fallout, which may spread and deposit over very large areas. If a nuclear yield is not achieved, the result would likely resemble an RDD in which fissile weapons material was utilized.</td>
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<tr>
<td>Incident</td>
<td>An occurrence, natural or manmade, that requires a response to protect life or property. Incidents can, for example, include major disasters, emergencies, terrorist attacks, terrorist threats, civil unrest, wildland and urban fires, floods, HazMat spills, nuclear accidents, aircraft accidents, earthquakes, hurricanes, tornadoes, tropical storms, tsunamis, war-related disasters, public health and medical emergencies, and other occurrences requiring an emergency response.</td>
</tr>
<tr>
<td>Incident Action Plan (IAP)</td>
<td>An oral or written plan containing general objectives reflecting the overall agency strategy for managing an incident. It may include the identification of operational resources and assignments and/or attachments that provide direction and important information for management of the incident throughout its lifecycle. The Planning Section Chief, in coordination with other Incident Command and General Staff members, is responsible for developing the IAP.</td>
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<tr>
<td>Incident Annexes</td>
<td>Describe the concepts of operations to address specific hazards requiring specialized application of the FDA EOP.</td>
</tr>
<tr>
<td>Incident Command Post (ICP)</td>
<td>The field location where the primary functions are performed. The ICP may be collocated with the Incident Base or other incident facilities.</td>
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</table>
### Incident Command System (ICS)
A standardized on-scene emergency management construct designed to provide for the adoption of an integrated organizational structure that reflects the complexity and demands of single or multiple incidents, without being hindered by jurisdictional boundaries. ICS is designed to enable effective incident management by integrating a combination of facilities, equipment, personnel, procedures, and communications operating within a common organizational structure. It is used for all kinds of emergencies and is applicable to small as well as large and complex incidents. ICS is used by various jurisdictions and functional agencies, both public and private, to organize incident management operations.

### Incident Commander (IC)
The individual responsible for all incident activities at the field level, including the development of strategies and tactics and the ordering and release of resources. The IC has overall authority and responsibility for conducting incident operations and is responsible for the management of all incident operations.

### Influenza
A serious disease caused by viruses that infect the respiratory tract.

### Investigational New Drug (IND) Application
The IND regulations are at 21 CFR Part 312. Current Federal law requires that a new drug be the subject of an approved marketing application before it is transported or distributed across State lines unless the drug is shipped for investigational uses under an IND application. During a new drug’s early preclinical development, the sponsor’s primary goal is to determine if the product is reasonably safe for initial use in humans and if the compound exhibits pharmacological activity that justifies commercial development. When a product is identified as a viable candidate for further development, the sponsor then focuses on collecting the data and information necessary to establish that the product will not expose humans to unreasonable risks when used in limited early-stage clinical studies.

FDA’s role in the development of a new drug begins when the drug’s sponsor (usually the manufacturer or potential marketer), having screened the new molecule for pharmacological activity and acute toxicity potential in animals, wants to test its diagnostic or therapeutic potential in humans. At that point, the molecule changes in legal status under the FD&C Act and becomes a new drug subject to specific requirements of the drug regulatory system.

The IND application must contain information in three broad areas:

- **Animal Pharmacology and Toxicology Studies.** Preclinical data to permit an assessment as to whether the product is reasonably safe for initial testing in humans. Also included is any previous experience with the drug in humans (often foreign use).
- **Manufacturing Information.** Information pertaining to the composition, manufacture, stability, and controls used for manufacturing the drug substance and the drug product. This information is assessed to ensure that the company can adequately produce and supply consistent batches of the drug.
- **Clinical Protocols and Investigator Information.** Detailed protocols for proposed clinical studies to assess whether the initial-phase trials will expose subjects to unnecessary risks and whether later stage trials are designed in a way to provide reliable evidence concerning effectiveness. Also, information on the qualifications of clinical investigators—professionals (generally physicians) who oversee the administration of the experimental compound—to assess whether they are qualified to fulfill their clinical trial duties. Finally, commitments to obtain informed consent from the research subjects, to obtain review of the study by an Institutional Review Board (IRB), and to adhere to the IND regulations.

Once the IND is submitted, the sponsor must wait 30 calendar days before initiating any clinical trials unless FDA grants a waiver. During this time, FDA has an opportunity to review the IND for safety to ensure that research subjects will not be subjected to unreasonable risk.

There are two general categories of INDs:

- **Commercial.** An IND for which the sponsor intends for the product to be commercialized at some point. The sponsor for a commercial IND is generally a corporate entity or, in some cases, the National Institutes of Health (NIH) or an academic institution.

- **Research (Noncommercial).** An IND for which a study is being conducted for research purposes rather than commercial development. These INDs are generally sponsored by an individual investigator or an academic institution.

Within the two IND categories, there are several IND types:

- **Standard IND,** while not a regulatory term, refers to an IND submitted as described under 21 CFR 312.23. Such an IND may be either a commercial or research IND.

- **Emergency IND** refers to when emergency use of an investigational drug is permitted by FDA based on telephone or other rapid communication request in some circumstances in which there is not time for submission of an IND application under the regulations. See 21 CFR 312.36. The term emergency IND is also, however, sometimes used to describe an IND that is specific to a particular patient and as to which use of the investigational drug is justified because alternative treatments are not available.
<table>
<thead>
<tr>
<th>Glossary Term</th>
<th>Description</th>
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<tr>
<td>Isolation</td>
<td>A state of separation between persons or groups to prevent the spread of disease. The first published recommendations for isolation precautions in U.S. hospitals appeared as early as 1877, when a handbook recommended placing patients with infectious diseases in separate facilities. Isolation measures can be undertaken in hospitals or homes, as well as in alternative facilities.</td>
</tr>
<tr>
<td>Joint Field Office (JFO)</td>
<td>The primary Federal incident management field structure. The JFO is a temporary Federal facility that provides a central location for the coordination of Federal and SLTT governments and private sector and nongovernmental organizations (NGOs) with primary responsibility for response and recovery. The JFO structure is organized, staffed, and managed in a manner consistent with NIMS principles and is led by a Unified Coordination Group. Although the JFO uses an ICS structure, the JFO does not manage on-scene operations. Instead, the JFO focuses on providing support to on-scene efforts and conducting broader support operations that may extend beyond the incident site.</td>
</tr>
<tr>
<td>Joint Information Center (JIC)</td>
<td>An interagency entity established to coordinate and disseminate information for the public and media concerning an incident. JICs may be established locally, regionally, or nationally depending on the size and magnitude of the incident.</td>
</tr>
<tr>
<td>Lead District</td>
<td>The FDA District Office with primary responsibility for addressing an incident. For the majority of incidents involving FDA-regulated products, the District in which the event is occurring (e.g., the physical location where people have been affected) will generally assume the lead District role.</td>
</tr>
<tr>
<td>Liaison Officer (LNO)</td>
<td>A member of the Command Staff responsible for coordinating with external Federal agencies (e.g., HHS, CDC, USDA, DHS, FBI, or DOS). LNOs serve as direct links between the AIC and their assigned agencies, and work as members of the external agencies’ EOCs, providing input on FDA policies, activities, concerns, resource availability, and other incident-related matters.</td>
</tr>
<tr>
<td>Logistics Section</td>
<td>Section responsible for providing facilities, services, and material support for the incident.</td>
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<tr>
<td>Major Disaster</td>
<td>Under the Stafford Act, any natural catastrophe (including any hurricane, tornado, storm, high water, wind-driven water, tidal wave, tsunami, earthquake, volcanic eruption, landslide, mudslide, snowstorm, or drought) or, regardless of cause, any fire, flood, or explosion in any part of the United States that, in the determination of the President, causes damage of sufficient severity and magnitude to warrant major disaster assistance under the Stafford Act to supplement the efforts and available resources of States, local governments, and disaster relief organizations in alleviating the damage, loss, hardship, or suffering caused thereby.</td>
</tr>
<tr>
<td>Memorandum of Understanding (MOU)</td>
<td>A formal agreement between FDA and another government agency (Federal, State, or local) or private institution, or an informal agreement with foreign governments or other foreign institutions.</td>
</tr>
<tr>
<td>Mission Assignment (MA)</td>
<td>A task given to an agency component to perform within a given operational period that is based on operational objectives defined in the IAP.</td>
</tr>
<tr>
<td>National Disaster Medical System (NDMS)</td>
<td>A federally coordinated system that augments the Nation’s medical response capability. The overall purpose of the NDMS is to supplement an integrated national medical response capability for assisting State and local authorities in dealing with the medical impacts of major peace-time disasters and to provide support to the military and the U.S. Department of Veterans Affairs (VA) medical systems in caring for casualties evacuated back to the United States from overseas armed conventional conflicts. The NRF uses the NDMS, as part of HHS/ASPR, under ESF #8 – Public Health and Medical Services, to support Federal agencies in the management and coordination of the Federal medical response to major emergencies and federally declared disaster, including:</td>
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<tr>
<td>National Disaster Recovery Framework (NDRF)</td>
<td>Serves as guide to enable effective recovery support to disaster-impacted SLTT jurisdictions. It provides a flexible structure that enables disaster recovery managers to operate in a unified and collaborative manner. It also focuses on how best to restore, redevelop, and revitalize the health, social, economic, natural, and environmental fabric of the community and build a more resilient Nation.</td>
</tr>
<tr>
<td><strong>National Incident Management System (NIMS)</strong></td>
<td>Provides a systematic, proactive approach to guide departments and agencies at all levels of government, NGOs, and the private sector to work seamlessly to prevent, protect against, respond to, recover from, and mitigate the effects of incidents, regardless of cause, size, location, or complexity, to reduce the loss of life and property and harm to the environment. NIMS works hand-in-hand with the NRF. It provides the template for the management of incidents, while the NRF provides the structure and mechanisms for national-level policy for incident management.</td>
</tr>
<tr>
<td><strong>National Infrastructure Protection Plan (NIPP)</strong></td>
<td>Plan that provides a coordinated approach to critical infrastructure and key resources (CIKR) protection for Federal, State, tribal, local, and private sector security partners. It sets national priorities, goals, and requirements for effective distribution of funding and resources that will help ensure that our government, economy, and public services continue in the event of a terrorist attack or other disaster.</td>
</tr>
<tr>
<td><strong>National Operations Center (NOC)</strong></td>
<td>Serves as the primary national hub for situational awareness and operations coordination across the Federal government for incident management. The NOC provides the Secretary of Homeland Security and other principals with information necessary to make critical national-level incident management decisions.</td>
</tr>
<tr>
<td><strong>National Preparedness Guidelines</strong></td>
<td>Guidance that establishes a vision for national preparedness and provides a systematic approach for prioritizing preparedness efforts across the Nation. These guidelines focus policy, planning, and investments at all levels of government and the private sector.</td>
</tr>
<tr>
<td><strong>National Response Framework (NRF)</strong></td>
<td>Guides how the Nation conducts all-hazards response. The NRF documents the key response principles, roles, and structures that organize national response. It describes how communities, States, the Federal government, and private sector and nongovernmental partners apply these principles for a coordinated, effective national response. Also, it details special circumstances where the Federal government exercises a larger role, including incidents where Federal interests are involved and catastrophic incidents where a State would require significant support. The NRF allows first responders, decision makers, and supporting entities to provide a unified national response.</td>
</tr>
<tr>
<td><strong>National Terrorism Advisory System (NTAS)</strong></td>
<td>The NTAS communicates information about terrorist threats by providing timely, detailed information to the public, government agencies, first responders, airports and other transportation hubs, and the private sector.</td>
</tr>
<tr>
<td><strong>New Drug Application (NDA)</strong></td>
<td>The NDA is the vehicle through which drug sponsors formally propose that FDA approve a new pharmaceutical for sale and marketing in the United States. The data gathered during the animal studies and human clinical trials of an IND become part of the NDA. The goals of the NDA are to provide enough information to permit the FDA reviewer to reach the following key decisions:</td>
</tr>
<tr>
<td><strong>Nuclide</strong></td>
<td>A general term applicable to all atomic forms of an element. The number of protons and neutrons in the nucleus and the amount of energy contained within the atom characterize nuclides.</td>
</tr>
<tr>
<td><strong>Offices</strong></td>
<td>Refers to the collective FDA headquarters and field offices, including those of the OC and ORA.</td>
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</tbody>
</table>
### Operating Division (OPDIV)
Agencies within HHS. HHS OPDIVs include FDA and the following:

- Administration for Children and Families (ACF)
- Administration on Aging (AoA)
- Agency for Healthcare Research and Quality (AHRQ)
- Agency for Toxic Substances and Disease Registry (ATSDR)
- Centers for Disease Control and Prevention (CDC)
- Centers for Medicare & Medicaid Services (CMS)
- Health Resources and Services Administration (HRSA)
- Indian Health Service (IHS)
- National Institutes of Health (NIH)
- Substance Abuse and Mental Health Services Administration (SAMHSA)

### Operations Section
The ICS Section responsible for all tactical incident operations and implementation of the IAP. In ICS, the Operations Section normally includes subordinate branches, divisions, and/or groups.

### Organizational Component
As defined by FDA Staff Manual Guide (SMG) 1005.1, “FDA Organizational Changes – Policy, Policy Regarding Organizational Changes,” any part of the FDA organization that is separately established as an organizational entity by law, regulation, the Secretary, or an official who has been delegated authority and that has formally assigned functions and an approved Standard Administrative Code and title. FDA organizational components include all Centers, Offices, divisions, branches, and resident posts at the headquarters and field levels.

### Pandemic
The worldwide outbreak of a disease in humans in numbers clearly in excess of normal. (See Epidemic.)

### Planning Section
The ICS Section responsible for the collection, evaluation, and dissemination of operational information related to the incident and for the preparation and documentation of the IAP. This Section also maintains information on the current and forecasted situation and on the status of resources assigned to the incident.

### Plume
The material spreading from a particular source and traveling through environmental media such as air or ground water. For example, a plume could describe the dispersal of particles, gases, vapors, and aerosols in the atmosphere or the movement of contamination through an aquifer (e.g., dilution, mixing, or adsorption onto soil).

### Preparedness
Actions that involve a combination of planning, resources, training, exercising, and organizing to build, sustain, and improve operational capabilities. Preparedness is the process of identifying the personnel, training, and equipment needed for a wide range of potential incidents and developing specific plans for the delivery of agency capabilities. While not part of the phases of FDA emergency operations, preparedness ensures that agency resources are available and effectively implemented when needed.

Directive for the development of a national preparedness goal and national preparedness system. PPD-8 is designed to facilitate an integrated all-of-Nation/whole community capabilities-based approach to preparedness.

### Presumptive
Used to describe information that strongly suggests a problem exists. Presumptive information consists of:

- Epidemiological data that provides a significant association between an illness, injury, or unanticipated adverse reaction and a product.
- An original analysis by a reliable laboratory revealing a significant level of a toxic substance in a regulated product, but confirmation is not complete.
- Natural or manmade incident reporting the extent of which is unknown at the present time.

### Prevention
Actions to avoid an incident or to intervene to stop an incident from occurring. It involves applying intelligence and other information to a range of activities that may include such countermeasures as deterrence operations, heightened inspections, investigations to determine the full nature and source of the threat, public health and agricultural surveillance and testing processes, and, as appropriate, specific law enforcement operations aimed at deterring, preempting, intimidating, or disrupting illegal activity and apprehending potential perpetrators and bringing them to justice.

### Program Executive Group
When there is an IMT, the PEG is established to provide program specific overarching strategy and policy guidance at the district/geographic level.

### Protection
Actions to mitigate the overall risk to FDA-regulated products resulting from accidental or intentional contamination/adulteration. Protection includes actions to deter the threat, mitigate vulnerabilities, or minimize consequences associated with a terrorist attack or other potential incident.
## Glossary of Terms

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Device (RDD)</strong></td>
<td>A device that disperses radioactive material by conventional explosive or other mechanical means, such as a spray.</td>
</tr>
<tr>
<td><strong>Program (SLEP)</strong></td>
<td>Pursuant to Section 319 of the PHSA, a determination by the HHS Secretary that a disease or disorder presents a public health risk, including significant outbreaks of infectious diseases or bioterrorist attacks. Existence of a public health emergency authorizes the Secretary to take certain actions appropriate to respond to the incident or potential incident, including making grants, providing awards for expenses, and entering into contracts and conducting and supporting investigations into the cause, treatment, or prevention of a disease or disorder.</td>
</tr>
<tr>
<td><strong>Shelf Life Extension</strong></td>
<td>DoD, the CDC, and VA maintain significant reserves of critical medical supplies. CDC maintains its reserves in the SNS. These supplies include drug products that have expiration dates. Constantly replacing drug products as they reach their expiration dates can be quite costly. To reduce overall costs to the Federal government and to the taxpayer, DoD, CDC, and VA work with FDA to determine whether a drug product’s useful life can be extended beyond the expiration date. FDA tests samples of specified lots of stored drug products submitted by participating Federal entities and analyzes the data to determine whether the data are adequate to allow for the use of product beyond the expiration date on the product labeling. The stored drug products are to be maintained under each product’s labeled storage conditions.</td>
</tr>
<tr>
<td><strong>Secretary</strong></td>
<td>Secretary of the U.S. Department of Health and Human Services (HHS).</td>
</tr>
<tr>
<td><strong>Response</strong></td>
<td>Immediate actions to protect consumers and ensure the safety of FDA-regulated products. Response also includes the execution of agency emergency plans and procedures and actions to support and mitigation of short-term recovery. Response activities may include applying intelligence and other information to lessen the effects or consequences of an incident; investigations into nature and source of the threat; ongoing public health and product surveillance and testing processes; and law enforcement operations aimed at preempting, interdicting, or disrupting illegal activity; and apprehending actual perpetrators and bringing them to justice.</td>
</tr>
<tr>
<td><strong>Resources</strong></td>
<td>Personnel and major items of agency equipment, supplies, and facilities available or potentially available for assignment to incident operations and for which status is maintained.</td>
</tr>
<tr>
<td><strong>Public Health</strong></td>
<td>The science and practice of protecting and improving the overall health of the community through disease prevention and early diagnosis, control of communicable diseases, health education, injury prevention, sanitation, and protection from environmental hazards.</td>
</tr>
<tr>
<td><strong>Public Health Emergency</strong></td>
<td>Pursuant to Section 319 of the PHSA, a determination by the HHS Secretary that a disease or disorder presents a public health risk, including significant outbreaks of infectious diseases or bioterrorist attacks. Existence of a public health emergency authorizes the Secretary to take certain actions appropriate to respond to the incident or potential incident, including making grants, providing awards for expenses, and entering into contracts and conducting and supporting investigations into the cause, treatment, or prevention of a disease or disorder.</td>
</tr>
<tr>
<td><strong>Public Information Officer (PIO)</strong></td>
<td>A member of the Command Staff responsible for managing incident-related information requirements. The PIO coordinates development and release of agency public health and consumer protection messages for use with the public, media, and/or other agencies with approval from FDA headquarters and HHS.</td>
</tr>
<tr>
<td><strong>Quarantine</strong></td>
<td>The period of isolation decreed to control the spread of disease. Before the era of antibiotics, quarantine was one of the few available means of halting the spread of infectious disease. It is still employed today as needed. The list of quarantinable diseases in the United States is established by Executive Order of the President, on recommendation of the HHS Secretary, and includes cholera, diphtheria, infectious tuberculosis, plague, smallpox, yellow fever, and viral hemorrhagic fevers (such as Marburg, Ebola, and Congo-Crimean disease). In 2003, severe acute respiratory syndrome (SARS) was added as a quarantinable disease. In 2005, another disease was added to the list, influenza caused by novel or reemergent influenza viruses that are causing, or have the potential to cause, a pandemic.</td>
</tr>
<tr>
<td><strong>Radiation</strong></td>
<td>Energy moving in the form of particles or waves. Familiar radiations are heat, light, radio waves, and microwaves. Ionizing radiation is a very high-energy form of electromagnetic radiation.</td>
</tr>
<tr>
<td><strong>Radioactive Contamination</strong></td>
<td>The deposition of unwanted radioactive material on the surfaces of structures, areas, objects, or people. It can be airborne, external, or internal.</td>
</tr>
<tr>
<td><strong>Radioisotope (Radioactive Isotope)</strong></td>
<td>Isotopes of an element that have an unstable nucleus. Science, industry, and medicine commonly use radioactive isotopes. The nucleus eventually reaches a stable number of protons and neutrons through one or more radioactive decays. Roughly 3,700 natural and artificial radioisotopes have been identified.</td>
</tr>
<tr>
<td><strong>Radiological Dispersal Device (RDD)</strong></td>
<td>A device that disperses radioactive material by conventional explosive or other mechanical means, such as a spray.</td>
</tr>
<tr>
<td><strong>Radionuclide</strong></td>
<td>An unstable and therefore radioactive form of a nuclide.</td>
</tr>
<tr>
<td><strong>Recovery (Short Term)</strong></td>
<td>The transition from emergency response back to normal operations. During the short-term recovery phase, FDA analyzes the cause of the incident to minimize the chance of the incident, emergency, or crisis reoccurring and repairs any strained relationships with stakeholders. The agency also assesses the actions taken to resolve the incident and addresses any short- or long-term effects. As a result, agency policies, plans, and procedures are updated, incorporating lessons learned from the event.</td>
</tr>
<tr>
<td><strong>Resources</strong></td>
<td>Personnel and major items of agency equipment, supplies, and facilities available or potentially available for assignment to incident operations and for which status is maintained.</td>
</tr>
</tbody>
</table>

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**FDA Emergency Operations Plan**
### Glossary of Terms

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shielding</td>
<td>The use of material placed between a radiation source and an individual to reduce exposure.</td>
</tr>
<tr>
<td>Sievert (Sv)</td>
<td>A unit of dose equivalent. This relates the absorbed dose in human tissue to the effective biological damage of the radiation. Not all radiation has the same biological effect, even for the same amount of absorbed dose. Dose equivalent is often expressed as millionths of a Sv, or microsievert (µSv). One Sv is equivalent to 100 roentgen equivalent man (rem).</td>
</tr>
<tr>
<td>Situation Report (SitRep)</td>
<td>A situation report or SitRep is a document that contains confirmed or verified information regarding the specific details relating to an incident. It provides explicit details (who, what, when, where, and how) of an incident.</td>
</tr>
<tr>
<td>Situational Awareness</td>
<td>The ability to identify, process, and comprehend the critical elements of information about an incident.</td>
</tr>
<tr>
<td>Stakeholder</td>
<td>Any person or entity involved in or affected by an FDA course of action. Stakeholders include consumers, FDA employees, vendors, and external government agencies.</td>
</tr>
<tr>
<td>Standard Operating Procedure (SOP)</td>
<td>A complete reference document or an operations manual that provides the purpose, authorities, duration, and details for the preferred method of performing a single function or a number of interrelated functions in a uniform manner.</td>
</tr>
<tr>
<td>Status Report</td>
<td>Information specifically related to the availability or assignment of agency resources. Status reports are used by Center, ORA Senior Emergency Response Coordinator, and/or District emergency coordinators to provide FDA senior officials with real-time information on the status of agency resources (i.e., personnel, facilities, equipment, and other materials) impacted by an incident and/or those deployed in support of emergency response operations. FDA status reports generally come in the form of daily teleconferences, or more frequently depending on situation requirements, sponsored by the Planning Section Chief.</td>
</tr>
<tr>
<td>Strategic National Stockpile (SNS)</td>
<td>The SNS is a national repository of vaccines, antivirals, antibiotics, chemical antidotes, antitoxins, life-support medications, intravenous administration, airway maintenance supplies, and medical/surgical items. The SNS is designed to supplement and resupply State and local public health agencies in the event of a national emergency anywhere and at anytime within the United States or its territories.</td>
</tr>
<tr>
<td>Surveillance and Detection</td>
<td>The ongoing, systematic collection, analysis, and interpretation of public health data essential to the planning, execution, and evaluation of FDA emergency operations closely integrated with the timely dissemination of these data to those responsible for prevention and control.</td>
</tr>
<tr>
<td>Technical Specialists</td>
<td>Person with special skills that can be used anywhere within the ICS organization. No minimum qualifications are prescribed, as Technical Specialists normally perform the same duties during an incident that they perform in their everyday jobs.</td>
</tr>
<tr>
<td>Terrorism</td>
<td>As defined under the Homeland Security Act of 2002, any activity that involves an act dangerous to human life or potentially destructive of CIKR; is a violation of the criminal laws of the United States or of any State or other subdivision of the United States in which it occurs; and is intended to intimidate or coerce the civilian population or influence or affect the conduct of a government by mass destruction, assassination, or kidnapping.</td>
</tr>
<tr>
<td>Threat</td>
<td>An indication of possible violence, harm, or danger.</td>
</tr>
<tr>
<td>Unified Command (UC)</td>
<td>An ICS application used when more than one agency has incident jurisdiction or when incidents cross political jurisdictions. Agencies work together through the designated members of the UC, often the senior persons from agencies and/or disciplines participating in the UC, to establish a common set of objectives and strategies and a single IAP.</td>
</tr>
<tr>
<td>Vaccine</td>
<td>A preparation consisting of antigens of a disease-causing organism that, when introduced into the body, stimulates the production of specific antibodies or altered cells. This produces immunity to the disease-causing organism. The antigen in the preparation can be whole disease-causing organisms (killed or weakened) or parts of these organisms.</td>
</tr>
<tr>
<td>Virus</td>
<td>Any of various simple submicroscopic parasites of plants, animals, and bacteria that often cause disease and that consist essentially of a core of ribonucleic acid (RNA) or deoxyribonucleic acid (DNA) surrounded by a protein coat. Unable to replicate without a host cell, viruses are typically not considered living organisms.</td>
</tr>
<tr>
<td>Warning Letter</td>
<td>A Warning Letter is a written communication from FDA notifying an individual or firm that the agency considers one or more products, practices, processes, or other activities to be in violation of the FD&amp;C Act or other relevant statutes, and that failure of the responsible party to take appropriate and prompt action to correct and prevent any future repeat of the violation may result in administrative and/ or regulatory enforcement action without further notice.</td>
</tr>
<tr>
<td>World Health Organization (WHO)</td>
<td>An agency of the United Nations established in 1948 to further international cooperation in improving health conditions.</td>
</tr>
</tbody>
</table>
This appendix provides the meaning of acronyms and abbreviations used in the FDA EOP.

<table>
<thead>
<tr>
<th>Acronym/Abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>µg/kg</td>
<td>Microgram Per Kilogram</td>
</tr>
<tr>
<td>µSv</td>
<td>Microsievert</td>
</tr>
<tr>
<td>24/7</td>
<td>24 Hours a Day, 7 Days a Week</td>
</tr>
<tr>
<td>A&amp;F</td>
<td>Administration and Finance [HHS/ASPR]</td>
</tr>
<tr>
<td>AABB</td>
<td>American Association of Blood Banks</td>
</tr>
<tr>
<td>AAR</td>
<td>After Action Report</td>
</tr>
<tr>
<td>ACCP</td>
<td>Assistant Commissioner for Compliance Policy</td>
</tr>
<tr>
<td>ACF</td>
<td>Administration for Children and Families [HHS]</td>
</tr>
<tr>
<td>ACO</td>
<td>Assistant Commissioner for Operations</td>
</tr>
<tr>
<td>ACOMS</td>
<td>Advisory Committee Oversight and Management Staff</td>
</tr>
<tr>
<td>ACRA</td>
<td>[Office of the] Associate Commissioner for Regulatory Affairs [ORA]</td>
</tr>
<tr>
<td>ADE</td>
<td>Adverse Drug Experience</td>
</tr>
<tr>
<td>Advisory Team</td>
<td>Advisory Team for Environment, Food, and Health</td>
</tr>
<tr>
<td>AEG</td>
<td>Agency Executive Group</td>
</tr>
<tr>
<td>AERS</td>
<td>Adverse Event Reporting System</td>
</tr>
<tr>
<td>AHRQ</td>
<td>Agency for Healthcare Research and Quality [HHS]</td>
</tr>
<tr>
<td>AIC</td>
<td>Agency Incident Coordinator</td>
</tr>
<tr>
<td>ALC</td>
<td>Agency Location Code</td>
</tr>
<tr>
<td>Am</td>
<td>Americium</td>
</tr>
<tr>
<td>AMS</td>
<td>Agricultural Marketing Service [USDA]</td>
</tr>
<tr>
<td>ANDA</td>
<td>Abbreviated New Drug Application</td>
</tr>
<tr>
<td>AoA</td>
<td>Administration on Aging [HHS]</td>
</tr>
<tr>
<td>ASL</td>
<td>[Office of the] Assistant Secretary for Legislation [HHS]</td>
</tr>
<tr>
<td>ASPA</td>
<td>[Office of the] Assistant Secretary for Public Affairs [HHS]</td>
</tr>
<tr>
<td>ASPR</td>
<td>[Office of the] Assistant Secretary for Preparedness and Response [HHS]</td>
</tr>
<tr>
<td>ATCC</td>
<td>American Type Culture Collection</td>
</tr>
<tr>
<td>ATS</td>
<td>Automated Targeting System [DHS/CBP]</td>
</tr>
<tr>
<td>ATSDR</td>
<td>Agency for Toxic Substances and Disease Registry [HHS]</td>
</tr>
<tr>
<td>BARDA</td>
<td>Biomedical Advanced Research and Development Authority</td>
</tr>
<tr>
<td>BASIS</td>
<td>Blood Availability and Safety Information System</td>
</tr>
<tr>
<td>BDPR</td>
<td>Biological Product Deviation Reporting</td>
</tr>
<tr>
<td>Bioterrorism Act</td>
<td>Public Health Security and Bioterrorism Preparedness and Response Act of 2002</td>
</tr>
<tr>
<td>BPDR</td>
<td>Biological Product Deviation Reporting</td>
</tr>
<tr>
<td>Bq</td>
<td>Becquerel</td>
</tr>
<tr>
<td>Bq/kg</td>
<td>Becquerel Per Kilogram</td>
</tr>
<tr>
<td>CAERS</td>
<td>CFSAN [Center for Food Safety and Applied Nutrition] Adverse Event Reporting System</td>
</tr>
<tr>
<td>CAN</td>
<td>Counterfeit Alert Network</td>
</tr>
<tr>
<td>CBER</td>
<td>Center for Biologics Evaluation and Research</td>
</tr>
<tr>
<td>CBP</td>
<td>U.S. Customs and Border Protection [DHS]</td>
</tr>
<tr>
<td>CBRN</td>
<td>Chemical, Biological, Radiological, and Nuclear</td>
</tr>
<tr>
<td>CBRNE</td>
<td>Chemical, Biological, Radiological, Nuclear, and High-Yield Explosive(s)</td>
</tr>
<tr>
<td>CC</td>
<td>Collaborating Centre [WHO]</td>
</tr>
<tr>
<td>Acronym</td>
<td>Full Form</td>
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<tr>
<td>---------</td>
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</tr>
<tr>
<td>CDC</td>
<td>Centers for Disease Control and Prevention</td>
</tr>
<tr>
<td>CDER</td>
<td>Center for Drug Evaluation and Research</td>
</tr>
<tr>
<td>CDRH</td>
<td>Center for Devices and Radiological Health</td>
</tr>
<tr>
<td>CEC</td>
<td>Center Emergency Coordinator</td>
</tr>
<tr>
<td>CFR</td>
<td>Code of Federal Regulations</td>
</tr>
<tr>
<td>CFSAN</td>
<td>Center for Food Safety and Applied Nutrition</td>
</tr>
<tr>
<td>cGMP or CGMP</td>
<td>Current Good Manufacturing Practice</td>
</tr>
<tr>
<td>CIKR</td>
<td>Critical Infrastructure and Key Resources</td>
</tr>
<tr>
<td>CMO</td>
<td>Chief Medical Officer</td>
</tr>
<tr>
<td>CMS</td>
<td>Centers for Medicare &amp; Medicaid Services [HHS]</td>
</tr>
<tr>
<td>COBOP</td>
<td>Continuity of Business Operations Plan</td>
</tr>
<tr>
<td>CONOPS</td>
<td>Concept of Operations</td>
</tr>
<tr>
<td>COOP</td>
<td>Continuity of Operations</td>
</tr>
<tr>
<td>CORE</td>
<td>Coordinated Outbreak Response and Evaluation</td>
</tr>
<tr>
<td>CPG</td>
<td>Comprehensive Preparedness Guide</td>
</tr>
<tr>
<td>CPGM</td>
<td>Compliance Program Guidance Manual</td>
</tr>
<tr>
<td>Cs</td>
<td>Cesium</td>
</tr>
<tr>
<td>CSR</td>
<td>CDER [Center for Drug Evaluation and Research] Situation Room</td>
</tr>
<tr>
<td>CTP</td>
<td>Center for Tobacco Products</td>
</tr>
<tr>
<td>CVM</td>
<td>Center for Veterinary Medicine</td>
</tr>
<tr>
<td>DACRA</td>
<td>Deputy Associate Commissioner for Regulatory Affairs</td>
</tr>
<tr>
<td>DD</td>
<td>District Director</td>
</tr>
<tr>
<td>DEOC</td>
<td>Director’s Emergency Operations Center</td>
</tr>
<tr>
<td>DERC</td>
<td>District Emergency Response Coordinator</td>
</tr>
<tr>
<td>DFAS</td>
<td>Defense Financial Accounting Center</td>
</tr>
<tr>
<td>DFDT</td>
<td>Division of Food Defense Targeting</td>
</tr>
<tr>
<td>DFI</td>
<td>Division of Field Investigations [ORA]</td>
</tr>
<tr>
<td>DFO</td>
<td>Division of Financial Operations [HHS/PSC]</td>
</tr>
<tr>
<td>DHS</td>
<td>U.S. Department of Homeland Security</td>
</tr>
<tr>
<td>DIB</td>
<td>Director, Investigations Branch</td>
</tr>
<tr>
<td>DIL</td>
<td>Derived Intervention Level</td>
</tr>
<tr>
<td>DIOP</td>
<td>Division of Import Operations and Policy</td>
</tr>
<tr>
<td>DMPTI</td>
<td>Division of Medical Products and Tobacco Inspections</td>
</tr>
<tr>
<td>DMPTPO</td>
<td>Division of Medical Products and Tobacco Program Operations</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic Acid</td>
</tr>
<tr>
<td>DoD</td>
<td>U.S. Department of Defense</td>
</tr>
<tr>
<td>DOE</td>
<td>U.S. Department of Energy</td>
</tr>
<tr>
<td>DOI</td>
<td>U.S. Department of the Interior</td>
</tr>
<tr>
<td>DOJ</td>
<td>U.S. Department of Justice</td>
</tr>
<tr>
<td>DOL</td>
<td>U.S. Department of Labor</td>
</tr>
<tr>
<td>DOS</td>
<td>U.S. Department of State</td>
</tr>
<tr>
<td>DUNS</td>
<td>Data Universal Numbering System</td>
</tr>
<tr>
<td>EIN</td>
<td>Employer Identification Number</td>
</tr>
<tr>
<td>eLEXNET</td>
<td>Electronic Laboratory Exchange Network</td>
</tr>
<tr>
<td>EMEA</td>
<td>European Medicines Agency</td>
</tr>
<tr>
<td>EMG</td>
<td>Emergency Management Group</td>
</tr>
<tr>
<td>EOC</td>
<td>Emergency Operations Center</td>
</tr>
<tr>
<td>EON-IMS</td>
<td>Emergency Operations Network – Incident Management System</td>
</tr>
<tr>
<td>EOP</td>
<td>Emergency Operations Plan</td>
</tr>
<tr>
<td>EPA</td>
<td>U.S. Environmental Protection Agency</td>
</tr>
<tr>
<td>Acronym</td>
<td>Description</td>
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<tr>
<td>----------</td>
<td>---------------------------------------------------------</td>
</tr>
<tr>
<td>EPEES</td>
<td>Emergency Planning, Exercises, and Evaluation Staff</td>
</tr>
<tr>
<td>Epi-X</td>
<td>Epidemic Information Exchange</td>
</tr>
<tr>
<td>ERCG</td>
<td>Emergency Response Coordination Group</td>
</tr>
<tr>
<td>ERIC</td>
<td>Employee Resource and Information Center</td>
</tr>
<tr>
<td>ERL</td>
<td>Essential Regulatory Laboratory [WHO]</td>
</tr>
<tr>
<td>ERP</td>
<td>Emergency Response Plan</td>
</tr>
<tr>
<td>ESEMS</td>
<td>Employee Safety and Environmental Management Staff [OO]</td>
</tr>
<tr>
<td>ESF</td>
<td>Emergency Support Function</td>
</tr>
<tr>
<td>EUA</td>
<td>Emergency Use Authorization</td>
</tr>
<tr>
<td>FACTS</td>
<td>Field Accomplishments and Compliance Tracking System</td>
</tr>
<tr>
<td>FATA</td>
<td>Federal Anti-Tampering Act</td>
</tr>
<tr>
<td>FBI</td>
<td>Federal Bureau of Investigation</td>
</tr>
<tr>
<td>FCC</td>
<td>Forensic Chemistry Center</td>
</tr>
<tr>
<td>FDA</td>
<td>U.S. Food and Drug Administration</td>
</tr>
<tr>
<td>FDRC</td>
<td>Federal Disaster Recovery Coordinator</td>
</tr>
<tr>
<td>FEMA</td>
<td>Federal Emergency Management Agency</td>
</tr>
<tr>
<td>FERN</td>
<td>Food Emergency Response Network</td>
</tr>
<tr>
<td>FOH</td>
<td>Federal Occupational Health</td>
</tr>
<tr>
<td>FOIA</td>
<td>Freedom of Information Act</td>
</tr>
<tr>
<td>FoodNet</td>
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U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES
U.S. Food and Drug Administration

Nuclear/Radiological Annex to the
FDA Emergency Operations Plan

Version 3.0

July 2019

OFFICE OF EMERGENCY MANAGEMENT
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## RECORD OF CHANGES

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A. **INTRODUCTION**

_Nuclear/Radiological emergencies_, whether accidental or intentional, have the potential to cause adverse health and safety effects for large segments of the human and animal populations. In order to mitigate the consequences of such incidents, the U.S. Food and Drug Administration (FDA) must possess the resources and capabilities necessary to prevent, prepare for, protect against, and rapidly and effectively respond to and recover from radiological incidents. A planned and coordinated approach to emergency operations by FDA organizational components in support of Federal, State, local, tribal, and territorial (SLTT) governments; international authorities; and responders in the field can save lives and ensure critical public health and medical needs are met.

This **Nuclear/Radiological Annex to the FDA Emergency Operations Plan (EOP)** establishes a framework for FDA’s management of radiological incidents. It provides the measures, operating structures, roles and responsibilities, and mechanisms for direction and coordination of FDA resources before, during, and after radiological emergencies that pose a risk to human or animal health. This Nuclear/Radiological Annex is compatible with the scalable, flexible, and adaptable Federal government emergency coordinating structures of the **National Response Framework (NRF)** and is consistent with the concepts, principles, and terminology of the National Incident Management System (NIMS)\(^60\) and the FDA EOP.\(^61\) This Nuclear/Radiological Annex is to be used, in conjunction with the FDA EOP, to assist FDA in conducting response operations for any radiological emergency.

A.1 **MISSION**

During a nuclear/radiological emergency, FDA must continue to meet its mission. In addition to the activities listed in the FDA EOP, FDA activities may expand to include performance of specific, incident-related functions such as:

- Activities conducted under the NRF’s **Nuclear/Radiological Incident Annex (NRIA)** and **Food and Agriculture Annex**, Emergency Support Functions (ESFs), and the NRF **Catastrophic Incident Supplement**.

- Assistance and recommendations to coordinating agencies and State, tribal, and local governments through the Advisory Team for the Environment, Food, and Health (Advisory Team).\(^62\)

A.2 **PURPOSE**

This Nuclear/Radiological Annex is intended to be used in conjunction with the FDA EOP and is not intended to be a “standalone” plan. The purpose of the Nuclear/Radiological Annex is to detail FDA responsibilities and operations specific to a radiological emergency and provide radiological-specific reference material to provide a coordinated and consistent agency approach to preparing for, preventing, protecting against, responding to, and recovering from radiological incidents involving or impacting FDA-regulated products. To accomplish this, the Nuclear/Radiological Annex:

- Is compatible with and expands on the FDA EOP, the overarching FDA-wide operational plan that addresses the full spectrum of natural and technological hazards and terrorist threats and the “umbrella plan” under which the Nuclear/Radiological Annex fits.

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\(^60\) For more information on the NRF and NIMS, refer to the “Authorities and References” section of this annex and FDA EOP Appendix A and Appendix B or visit [www.fema.gov/emergency/nrf/](http://www.fema.gov/emergency/nrf/).

\(^61\) For more information on how FDA responds to emergencies, refer to the FDA EOP.

\(^62\) For more information on the Advisory Team, refer to Appendix B of this Nuclear/Radiological Annex.
• Defines the FDA radiological emergency operations structure and assigns essential tasks to the 
FDA organizational components involved in prevention, protection, response, and recovery 
efforts.

A.3 SCOPE AND APPLICABILITY
The Nuclear/Radiological Annex covers the requirements in anticipation of or in response to a radiological or nuclear incident that affects or involves an FDA-regulated product. This annex applies to both inadvertent and deliberate nuclear/radiological events. According to the NRIA to the NRF, inadvertent releases include: incidents at nuclear facilities (commercial or weapons production facilities), lost radioactive material sources, transportation accidents involving nuclear/radioactive material, domestic nuclear weapons accidents, and foreign accidents involving nuclear or radioactive material that impact the United States or its territories, possessions, or territorial waters, and assistance to other nations and U.S. citizens abroad as requested. In addition, this annex includes, but is not limited to, response to the effects of deliberate attacks perpetrated with radiological dispersal devices (RDDs), nuclear weapons, or improvised nuclear devices (INDs). Refer to FDA EOP Appendix K, “Glossary of Terms” for RDD and IND definitions.

A.4 PLANNING ASSUMPTIONS
The Nuclear/Radiological Annex is based on the same planning assumptions as referenced in the FDA EOP and, in addition:

• The U.S. Department of Homeland Security (DHS) will review the situation and determine whether to assume Federal leadership for the overall response in accordance with the NRF.

• To ensure the capability to implement the Nuclear/Radiological Annex, each FDA headquarters and field organizational component tasked with radiological emergency roles and responsibilities, as identified in this Nuclear/Radiological Annex or the FDA EOP, may develop and maintain individual emergency plans and procedures that identify the critical and time-sensitive missions, functions, assignments, and processes to be performed by their organizational component during a radiological incident. These documents shall be consistent with the FDA EOP and this annex and available to all Center/Office personnel.

A.5 ACTIVATION
The ESF #8 – Public Health and Medical Services response to a radiological event will be directed by U.S. Department of Health and Human Services (HHS) Office of the Secretary, in coordination with DHS, which is responsible for managing the U.S. government response. FDA is a subordinate Operating Division (OPDIV) of HHS and will align its response activities with those of HHS. FDA will use the full spectrum of its resources to accomplish assigned roles, responsibilities, functions, goals, and missions.

This annex will be consulted and implemented together with the FDA EOP when there is a radiological incident. Examples of events that could trigger activation of this annex include, but are not limited to, any radiological incident, including nuclear power plant accidents, manmade accidents, RDDs, or INDs. Such events may impact FDA-regulated products/industries, lead to a response from FDA via the Advisory Team, or result in the request for the issuance/use of Emergency Use Authorizations (EUAs) or Emergency Investigational New Drug applications (EINDs) for a medical countermeasure.

District Directors (DDs), Program Directors (PDs) and Center/Office Directors may activate their individual organizational component’s EOP without activation of the FDA Nuclear/ Radiological Annex.
A.6 SUPERSEDERENCE

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B. CONCEPT OF OPERATIONS

The following concept of operations (CONOPS) describes the principal authorities’ governing agency emergency functions and the phases within which FDA conducts radiological incident-related operations.

B.1 EMERGENCY AUTHORITIES

FDA’s emergency authorities are prescribed by several statutes and conducted in accordance with applicable regulations as described in FDA EOP Section F, “Authorities and References.” These include the Pandemic and All-Hazards Preparedness Reauthorization Act (Public Law 113-5) and Title 21, “Food and Drugs,” of the Code of Federal Regulations (CFR). In addition to the authorities listed in the FDA EOP, the emergency authority relevant to a radiological/nuclear incident is:


B.2 EMERGENCY OPERATIONS PHASES

The following phases comprise the entire spectrum of FDA emergency operations: Prevention, Protection, Mitigation, Response, and Recovery. During a nuclear/radiological event FDA employees are expected to follow the activities described in the FDA EOP, Section B.2, “Emergency Operations Phases.” Although emergency operations may involve each of these phases over the course of a radiological incident, the nature and severity of an event and the FDA organizational component(s) responding will determine the specific order, actions, and responsible parties required for each. The following sections describe additional resources available during FDA’s emergency operations for prevention and protection specific to a radiological or nuclear event.

B.2.1 Prevention, Protection, and Mitigation

FDA participates in a number of planning subcommittees under the Federal Radiological Preparedness Coordinating Committee (FRPCC), which coordinates all Federal responsibilities for assisting State and local governments in their emergency planning and preparedness for peacetime radiological emergencies, developing protective action recommendations (PARs) for human and animal feed, providing guidance on radioprotective substances and prophylactic drugs to reduce radiation dose, and providing medical and mental health support through ESF #8. The Federal Emergency Management Agency (FEMA) is the chair of the FRPCC, and FDA serves as the chair of the FRPCC Subcommittee on Food, Health, and Environment.

The FRPCC performs the following functions:

- Assisting the FEMA Director in providing policy direction with respect to Federal assistance to State and local governments in their radiological emergency planning and preparedness activities.
- Establishing subcommittees to aid in carrying out its functions; current subcommittees include Training, Offsite Instrumentation, Transportation, and Federal Response.
- Assisting FEMA in resolving issues relating to granting final approval (under 44 CFR Part 350) of a State radiological emergency preparedness plan.

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63 A more detailed summary of the laws, regulations, directives, and policy guiding FDA emergency operations is provided in Section F, “Authorities and References,” of this annex and the FDA EOP.

64 Refer to Section C, “Organization and Assignment of Responsibilities,” of this annex for an overview of FDA organizational component roles and responsibilities during a radiological incident.
• Coordinating research and study efforts of its member agencies relative to State and local government radiological emergency preparedness to ensure minimum duplication and maximum benefits to State and local governments.

The FRPCC works directly with HHS, the Centers for Disease Control and Prevention (CDC), and FDA seeking guidance for State and local governments on the use of radioprotective substances and prophylactic use of drugs (e.g., potassium iodide, Radiogardase®) to reduce the radiation dose to specific organs, including dosage and projected radiation exposures.

B.2.1.1 Surveillance and Detection

B.2.1.1.1 Winchester Engineering and Analytical Center

Winchester Engineering and Analytical Center (WEAC) is an FDA/Office of Regulatory Affairs (ORA) laboratory that is equipped and staffed to handle the analysis of radioactive materials. It is the Federal laboratory having the greatest capability in the United States to analyze food samples for radioactive contamination. As such, WEAC serves as the lead for the radiological component of the Food Emergency Response Network (FERN).65 The FDA FERN program is administered by the ORA Division of Regulatory Science. FDA conducts import and domestic food surveillance programs for radioactive contamination (e.g., import samples, onsite authorities or SLTT investigator-provided samples, market basket survey samples).

B.2.2 Response

B.2.2.1 Gain and Maintain Situational Awareness

B.2.2.1.1 Alert and Notification

FDA may be alerted to a threat, hazard, or other significant event through a variety of means, including the surveillance systems discussed in the FDA EOP Table B-1 (Surveillance and Detection Systems Utilized by FDA) and/or directly from external Federal agencies (e.g., the HHS Secretary’s Operations Center [SOC], U.S. Environmental Protection Agency [EPA], DHS, U.S. Department of Energy [DOE], U.S. Nuclear Regulatory Commission [NRC], and/or the Federal Bureau of Investigation [FBI]); SLTT and international public health agencies; industry; consumers; news media; and internal FDA organizational components.

If FDA becomes aware of a radiological incident from a non-DHS agency, it should notify the coordinating agency and the DHS National Operations Center (NOC) at 202-282-8101 and comply with other appropriate statutory requirements for notification. If FDA becomes aware of an overt threat or act involving nuclear/radiological material/device or indications the event is not inadvertent or otherwise accidental, the U.S. Department of Justice (DOJ) should be notified through the FBI.

B.2.2.1.2 Assessment and Monitoring

During a radiological response, WEAC is the lead FDA laboratory for the analysis of radionuclides in foods, drugs, and medical devices for radiological emergency consequence management.

B.2.2.2 Medical Countermeasures

Information on available medical countermeasures and evaluation of EUA requests can be provided by the Office of Counterterrorism and Emerging Threats (OCET) or Center emergency coordinators, as needed, through their contact and work with Center subject matter experts (SMEs).

65 For more information on the radiological FERN and WEAC, refer to Appendix A of this Nuclear/Radiological Annex.
B.2.2.3 Activation and Deployment of Resources and Capabilities

FDA is a member and currently the chair of the Advisory Team, a Federal interagency group of radiation SMEs who provide coordinated, science-based advice and recommendations to the coordinating agency\(^66\) and/or the State, tribal, and local government on protective measures that may be taken following a radiological incident. The Advisory Team comprises representatives from the EPA, U.S. Department of Agriculture (USDA), and HHS, including the CDC and FDA. Other departments or agencies may be asked to participate as needed. The Advisory Team works closely with the Federal Radiological Monitoring and Assessment Center (FRMAC), an interagency group led by the DOE National Nuclear Security Agency (NNSA).\(^67\)

If Federal assistance is requested for a radiological/nuclear incident and Advisory Team participation is needed, FDA will send an ORA Radiological Health Representative(s) (RHR) to the Unified Command (UC) as part of the onsite Advisory Team. (See Appendix B for activation/notification procedures of the Advisory Team.) Other Advisory Team members will be available at headquarters or in other locations to provide reachback and coverage when the remote Advisory Team member(s) is in transit or otherwise unavailable.

B.2.2.4 Coordination of Response Actions

During the response phase, FDA’s role is to provide information about countermeasures and facilitate their availability as stated in the FDA EOP, Section B.2.2.3.5, “Medical Countermeasures: EUA and Expanded Access.” Through the Advisory Team, FDA provides guidance to State and local agencies on interdicting suspected food from the food supply, as well as support for making protective action decisions.

To assist FDA’s Emergency Operations Center (EOC) in responding to a radiological emergency, staff personnel in the medical product Center maintain the status of medical countermeasure product availability and relevant manufacturing information. Each Center has detailed policies and procedures in place to make critical medical countermeasures available, and the FDA EOC coordinates the sharing of this information during an emergency.

B.2.2.4.1 Exposure Determination Information

Based on data provided by FRMAC and other field organizations, the coordinating agency will provide the following information, which may be helpful in guiding decisions about medical countermeasures or other protective actions:

- Identity of the radioisotopes.
- Quality of radiation (alpha, beta, gamma, and neutrons).
- Nature of the incident (e.g., nuclear power plant release, IND or RDD).
- Casualty information, including estimates of combined radiation and non-radiation injuries, such as blast and burn.

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\(^66\) Coordinating agencies are those Federal agencies that own, have custody of, authorize, regulate, or are otherwise assigned responsibilities for the nuclear/radioactive material, facility, or activity involved in the incident. A listing of coordinating agencies can be found in Table 1 of the NRF NRIA. [www.fema.gov/pdf/emergency/nrf/nrf_nuclearradiologicalincidentannex.pdf](www.fema.gov/pdf/emergency/nrf/nrf_nuclearradiologicalincidentannex.pdf)

\(^67\) For additional information, see the NRF NIRA.
• Plume direction and characteristics (estimated population exposure), including Shape files, which can be used to generate geographic information system (GIS) maps for situational awareness and planning.

• Amount of external radiation exposure.

This information may assist coordinating agencies in predicting which people exposed to radiation may develop:

• Hematopoietic syndrome (projected needs for cytokines and antimicrobials).
• Gastrointestinal syndrome (possible use of investigational cytokines).
• Cardiac/central nervous system syndrome (following extremely high doses, with death likely).

**B.2.3 Recovery**

During the recovery phase, some FDA objectives that would support Federal or SLTT agencies are:

• To manage a potentially large number of samples of FDA-regulated products associated with nuclear/radiological releases, including long-term (months to years) sampling, product destructions, and product reconditioning, as applicable.

• To provide technical support to the Advisory Team in establishing “clean-up” levels in following a radiological incident.
C. ORGANIZATION AND ASSIGNMENT OF RESPONSIBILITIES

This section identifies the roles and responsibilities specific to a radiological incident of FDA headquarters and field organizational components in preventing and protecting against, responding to, and recovering from a radiological incident. FDA staffs work closely with one another and with governmental and industry partners to ensure the safety, efficacy, and security of FDA-regulated products that mitigate the public health effects of an emergency or disaster in the United States or worldwide. This is done by using novel and expeditious approaches to product regulation in order to optimize the availability and use of FDA-regulated products in all populations.

The following is a brief description of the radiological emergency functions performed by responsible organizational components. Refer to individual Center/Office emergency plans and procedural documentation, as applicable, for more detailed information on radiological emergency activities. For additional information on general FDA Center/Office emergency roles and responsibilities, refer to the FDA EOP, Section C, “Organization and Assignment of Responsibilities.”

FDA product Centers are responsible for the regulation of a defined set of products. Their professional staff includes both clinical and scientific experts. This expertise (e.g., analytical, laboratory, sampling procedures, subject matter expertise, and industry knowledge) is available for critical consultation in the event of a nuclear/radiological incident. In addition, the Center for Food Safety and Applied Nutrition (CFSAN) has responsibilities that are specific to a radiological incident.

C.1 OFFICE OF THE COMMISSIONER

C.1.1 Office of Operations

C.1.1.1 Office of Security and Emergency Management

In addition to the activities specified in the FDA EOP, the Office of Emergency Management (OEM) and Office of Emergency Operations (OEO) will communicate with external coordinating agencies and share information with internal FDA entities for nuclear/radiological incidents affecting FDA-regulated products. Dependent on the type of incident, some of the agencies with which FDA would communicate are: the U.S. Department of Defense (DoD), DOE, National Aeronautics and Space Administration (NASA), DHS, EPA, and the NRC. For a detailed description of what the NRF provides about the response and coordinating responsibilities, see the NRF NRIA at www.fema.gov/pdf/emergency/nrf/nrf_nuclearradiologicalincidentannex.pdf.

68 Those FDA organizational components without defined radiological emergency roles or responsibilities are not described within this annex.

The external coordinating agencies will share with OEM/OEO information about response activities of interest to FDA. The information to be shared with FDA includes but is not limited to:

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<td>Population Modeling Data</td>
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OEM/OEO will also be responsible for alert and notification of the Advisory Team in response to activation or consultation requests from Federal agencies and/or SLTTs. OEM/OEO will support Advisory Team operations by providing conference call support and interface with the FDA Incident Management Group (IMG), when activated.

**C.1.2 Office of Laboratory Science and Safety**

**C.1.2.1 Employee Safety and Environmental Management Staff**

The Employee Safety and Environmental Management (ESEM) Staff serves as the “corporate” ESH office for the Office of the Commissioner (OC), while each Center/Office is responsible for implementing operational ESH programs and does so with Center/Office ESH Managers. The ESEM Staff perform the enterprise functions of serving as principal advisors to FDA in all matters pertaining to safety and occupational health management, and is responsible for agency-wide safety policy development and overall program management, including recommendations on funding and training programs. The office also provides technical assistance to the Centers and field activities and is responsible for evaluating the safety and health programs throughout the agency through review and inspection. The ESEM staff provides official FDA safety representation to other safety, health and environmental-related regulatory agencies as well as Health and Human Services (HHS) and the General Services Administration (GSA).

Further, ESEM staff provide the operational enterprise functions of occupational medical oversight of the agency’s program; coordinating the agency’s Occupant Emergency preparedness, response and fire safety, establishing and administering a headquarters-wide Institutional Biosafety Committee (IBC); managing a comprehensive radiation safety program for the White Oak Campus including the administration of a single broad scope Nuclear Regulatory Commission (NRC) License; coordinating the hazardous waste disposal contract for the HQ area Centers; and project management, including risk management, to minimize agency liabilities when decommissioning laboratory space. Oversight and management of Campus waste water permit, hazardous waste permits and Emergency Planning and Community Right to Know Act (EPCRA) reporting.

**C.1.3 Office of Chief Scientist**

**C.1.3.1 Office of Counterterrorism and Emerging Threats**

OCET works with FDA’s medical product Centers to ensure the availability of medical countermeasures that can improve survival and mitigate and treat injuries from radiological/nuclear events and their delayed effects including acute radiation syndrome. These include decorption or blocking agents to reduce internal contamination from exposure to radionuclides and biodosimetry devices that can be used in a radiological/nuclear event to assess exposure to radiation and facilitate response activities.
C.2. Office of Foods and Veterinary Medicine

C.2.1 Center for Food Safety and Applied Nutrition

CFSAN, in conjunction with the agency’s field staff, is responsible for promoting and protecting the public’s health by ensuring that the Nation’s food supply is safe, sanitary, wholesome, and honestly labeled and that cosmetic products are safe and labeled properly. In addition to the activities described in the FDA EOP, CFSAN is responsible for, but not limited to, the following activities during radiological emergency:

- Overseeing the safety of the U.S. food supply following radiological emergencies through its Compliance Program and its participation with the Advisory Team, using the FDA food protective action guidance as the principal source of doctrine.
- Through its involvement with FRMAC, participates in the development of assessment methods and guidance for food protective measures.
- Providing scientific, technical, and policy expertise, guidance, and support within FDA and to other Federal and State agency components, regulated industry, and the public.
- Issuing recommendations and guidelines on acceptable levels of radioactive contamination of foods, dietary supplements, and cosmetics.
- Providing advice regarding health risks from radioactive contamination of food, dietary supplements, and cosmetics; this includes advice on actions that can be taken by FDA, State and local authorities, industry, or consumers to reduce public exposure to contaminated food, dietary supplements, and cosmetics.
- Overseeing the overall response to verify that the necessary food- and cosmetic-related actions are taken.
- Providing advice regarding sampling procedures and testing methodology (screening and confirmatory).
- Providing advice regarding product traceback and traceforward investigations.
- Developing field programs to meet intermediate to long-range consequence management needs.

C.3 Office of Global Regulatory Operations and Policy

C.3.1 Office of Regulatory Affairs

ORA, led by the Associate Commissioner for Regulatory Affairs (ACRA), protects consumers and enhances public health by maximizing compliance of FDA-regulated products and minimizing risk associated with those products. ORA is the lead office for all agency field activities. In addition to the activities described in FDA EOP, ORA is responsible for providing RHRs and other qualified ORA staff, as needed, to be FDA representatives to the field component of the Advisory Team.

C.3.1.1 Radiological Health Representatives

The RHRs represent the agency on committees chaired by FEMA that include representatives from the U.S. Public Health Service (USPHS), EPA, USDA, NRC, U.S. Coast Guard (USCG), and others. These

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70 The duties and responsibilities of the RRHRs are defined in Compliance Program Guidance Manual (CPGM) Program No. 7386.009, “Emergency Planning and Response Activities.”
committees are responsible for evaluating State and local emergency plans for nuclear power plants and radiological emergencies.

In addition to serving as representatives on these committees, the RHRs represent FDA on the onsite Advisory Team (Section B.2.2.3) during a radiological incident. Other FDA SMEs may also represent FDA on the Advisory Team through deployment or via telephone/email in a reachback status. FDA staff is activated for the Advisory Team as per the Advisory Team Activation Standard Operating Procedure (SOP) (see Appendix B). When an incident has occurred, RHRs are the agency’s onsite representatives to the Advisory Team. If there is a request for PARs for FDA-regulated products, RHRs help develop the recommendations, in consultation with other FDA scientific and policy experts. For additional information on PARs and Protective Action Guides (PAGs), refer to Appendix C, “Protective Actions.”
D. DIRECTION, CONTROL, AND COORDINATION

This section provides a description of radiological-specific positions and teams and describes their primary roles and responsibilities. These individuals are expected to have read and be familiar with the NRF NRIA (see www.fema.gov/pdf/emergency/nrf/nrf_nuclearradiologicalincidentannex.pdf).

FDA organizes emergency response operations in accordance with the concepts, principles, and terminology of the Incident Command System (ICS), as defined within NIMS. Incident Command and subordinate Planning, Operations, Logistics, and Finance/Administration functions lay the foundation for the agency’s implementation of ICS during all hazards, at headquarters and field levels, and across geographic divides. The inherent design of this system enables rapid, scalable, and flexible agency-wide emergency management activities.

D.1 FIELD INCIDENT COMMAND

FDA field response will be established as described in FDA EOP, Section B.2.2.3.2, “FDA Field Response” and Section D.1, “Field Incident Command.” Depending on the location of the incident and extent of potential contamination, an Incident Management Team (IMT) may be established. The IMT will coordinate with IMG, Advisory Team, and SLTT agencies to respond to an incident. FDA does not have a role in collecting data in restricted areas with high levels of radiological contamination.

D.2 HEADQUARTERS INCIDENT COORDINATION

As FDA’s coordination point for emergencies, due to the severity or potential consequences of a nuclear/radiological incident, an IMG will be established to address the public health impact. Figure D-1 depicts a notional IMG emergency response coordinating structure for a radiological incident.

![Figure D-1. Notional FDA IMG for a Nuclear/Radiological Event](image)

71 For more information on the ICS, refer to www.training.fema.gov/EMIWeb/IS/ICSResource/index.htm.
**D.2.1 Command Staff**

The FDA Command Staff includes a Public Information Officer (PIO), Liaison Officers (LNOs), and a Safety Officer.

**D.2.1.1 Radiological-Specific Liaisons**

When activated, the FDA EOC serves as the agency-wide focal point for radiological emergency operations, coordination, and communications for incidents involving regulated products. External liaisons from the USDA, NRC, or other agencies may be present at the FDA EOC. FDA may also send liaisons to the coordinating agency for the incident, or participate in the response coordination, if an Incident Command or UC is established.

**D.2.2 General Staff**

The General Staff is responsible for the functional aspects of the FDA ICS. The General Staff consists of Operations, Planning, Logistics, and Finance/Administration Sections composed of designated FDA headquarters and field emergency staff. Individual position assignments within these Sections are outlined in the FDA EOP.

**D.2.2.1 Operations Section**

The Operations Section is responsible for coordinating all agency field activities in response to a radiological/nuclear event involving a regulated product. Figure D-2 depicts FDA’s Operations Section during the event. The groups included in the chart below could possibly be established when the number of resources exceeds the manageable span of control of the IMG and Operations Section Chief. Groups are established to divide the incident into functional areas of operation. Additional levels of supervision may also exist below the group level.

![Operations Section Organizational Elements](image)

**Figure D-2. Notional Operations Section Organizational Elements**

Expansion or contraction of the Operations Section may vary according to numerous considerations and operational factors associated with an incident. In some cases, a functional approach may be used. In other cases, the organizational structure will be determined by geographical or Center/Office jurisdictional boundaries. In still others, a mix of functional and geographical considerations may be appropriate. The Agency Incident Coordinator (AIC) will determine the appropriate structural approach based on the specific circumstances of the incident at-hand.

**D.2.2.1.1 Drug Group**

The Drug Group addresses drug issues such as adverse event reporting, labeling changes, whether EUAs for drugs are appropriate, and what restrictions should be placed on uses authorized under EUAs related to a radiological operational response. Some potential teams, led by the Center for Drug Evaluation and Research (CDER) staff, that may be needed to manage mitigation and recovery issues include:
• Expired Drugs Team
• Drug Shortage Team
• Adverse Drug Events Team

**Expired Drugs Team**

The Expired Drugs Team deals with such issues as use of expired drugs and drugs out of specification, including those in the Strategic National Stockpile (SNS). This team would have input in discussing whether an EUA should/can include private sector stockpiles (e.g., hospital associations, large employers) or whether a separate authorization is needed. The Expired Drugs Team would also monitor labeling and Shelf Life Extension Program (SLEP) issues related to the event.

**Drug Shortage Team**

The Drug Shortage Team is responsible for reporting on the availability and surge capacity for drugs, monitoring commercial inventory, and working with manufacturers of drug products to prepare for possible increases in demand. The team would contact intravenous fluid manufacturers and other critical care drug manufacturers to prepare for possible increased needs.

**Adverse Drug Events Team**

The Adverse Drug Events Team assists with the monitoring of adverse drug events to identify undesired side effects and unexpected adverse effects from a drug product used for treatment.

**D.2.2.1.2 Biologics Group**

The Biologics Group addresses blood and other biologics issues as part of a radiological/nuclear operational response.

**D.2.2.1.3 Medical Device Group**

The Medical Device Group would manage issues related to a radiological/nuclear operational response. Potential teams, with leaders from Center for Devices and Radiological Health (CDRH) staff members, who may be needed to manage mitigation and recovery issues, are:

• In-Vitro Diagnostics (IVD) Team
• Personal Protective Equipment (PPE) and Other Devices Team
• Shortage Team

**In-Vitro Diagnostics Team**

The IVD Team addresses issues related to the following:

• Triage of EUA requests.
• Response to inquiries regarding EUA requests from other Federal agencies, and domestic and international commercial sources for IVDs claiming to assess radiation exposure.
• Contact with manufacturers regarding assays and instruments to be used in development of IVDs.
• Clear or approve tests and instruments that detect and measure in situ radioisotope contamination and absorbed radiation dose in exposed individuals.
• Monitor reagent and system shortages on a short-term basis (daily or weekly) as well as a long-term basis (monthly).
• Monitor radiation biodosimetry and in situ radioactivity contamination detection device functional claims.
• Review data and labeling of cleared or approved devices for which a company is proposing new
claims and changes.
• Address fraudulent claims by monitoring websites and complaints regarding product claims and
issue warning letters as appropriate.

**Personal Protective Equipment and Other Devices Team**

The PPE and Other Devices Team oversee or manage the following:

• Exchange of information with the CDC/SNS, States, and manufacturing sector and real-time
monitoring of consumption/utilization rates of PPE during emergencies.
• Respond to State and private sector inquiries about perceived shortages during public health
emergencies.
• Anticipate and plan for future surges in demand for devices.
• Track/trace status of devices released from SNS.
• Meet with manufacturers about the development of new PPE.
• Address PPE shortages by working with industry to update current manufacturing capabilities and
assist with long-term planning by the manufacturing sector to meet demands for potential surges.

**Shortage Team**

The Shortage Team would be created if shortages of IVD, PPE, and other devices are too extensive to be
managed by the above teams alone.

**D.2.2.1.4 Food/Feed Safety Group**

The Food/Feed Safety Group addresses issues related to the following:

• Providing scientific, technical, and policy expertise, guidance, and support within FDA and to
other Federal and State agency components, regulated industry, and the public.
• Providing advice regarding health risks from radioactive contamination of food and cosmetics
and advice on actions that can be taken by FDA, State and local authorities, industry, or
consumers to reduce public exposure to radiation.
• Evaluating and providing guidance for food- and cosmetic-related response actions.
• Advice regarding sampling plans and procedures and testing methods (screening and
confirmatory).
• Advice regarding product traceback and traceforward investigations.
• Developing field programs to meet intermediate to long-range consequence management needs.
D.2.2.2 Planning Section

The Planning Section is responsible for collecting, evaluating, and disseminating operational information pertaining to the incident. This Section maintains information and intelligence on the current and forecasted situation, as well as the status of FDA resources assigned to the incident. Technical Specialists may be assigned to the Planning Section or to other IMG Sections as determined by the AIC.

D.2.2.2.1 Radiological Emergency Response Unit

The Radiological Emergency Response Unit consists of a cadre of radiological health experts from relevant Centers/Offices with a variety of backgrounds, including health physics, medical physics, radiology, nuclear medicine, and radiation oncology. The unit provides technical advice to FDA senior management and liaisons to external agencies. The Unit Leader is identified by OEM in coordination with the FDA representative on the Advisory Team.

D.2.2.2.2 Technical Specialists

Technical Specialists such as Science/Medical Specialists and EUA/Product Interface Specialists are normally assigned to the Planning Section, but may support any ICS function including the Command Staff at the discretion of the AIC. 72

D.2.2.2.3 Employee Health Specialist

The Employee Health Specialist is a staff member of OCET or the Office of Operations (OO)/ESEMS who addresses the need for appropriate health protection for high-risk FDA employees. The Employee Health Specialist works with HHS and the USPHS Federal Occupational Health (FOH) Services to ensure adequate medical countermeasures are made available to protect FDA responders. The Employee Health Specialist is assigned to the Planning Section or can be called to serve in the Logistics Section.

72 See FDA EOP, Section D, “Direction, Control, and Coordination,” for a description of the Technical Specialist positions.
E. COMMUNICATIONS AND INFORMATION MANAGEMENT

Effective emergency management and incident response activities rely upon flexible communications and information systems that provide for a “common operating picture” across all agency components and staffs. Radiological/nuclear incident responders are expected to follow the FDA EOP, Section E, “Communications and Information Management.”

E.1 EMERGENCY ALERTS

FDA may receive alert and notification of a domestic nuclear/radiological incident from multiple external agencies. These notifications in the process may occur simultaneously following detection of an incident (see Figure E-1). Possible reporting pathways include alerts from:

A. FRMAC to the Advisory Team through the FDA EOC (see Appendix B). Because of existing protocols outlined in the NRIA-NRF, the FRMAC may initially contact the CDC Director’s Emergency Operations Center (DEOC), who will contact the FDA EOC for notification of the Advisory Team.

B. State or local government representative(s) to the Advisory Team through the FDA EOC. Because of existing protocols outlined in the NRIA-NRF, State and local government officials may initially contact the CDC DEOC, who will contact the FDA EOC for notification of the Advisory Team.

C. NRC to the Advisory Team through the FDA EOC.

D. NRC to FEMA to the HHS SOC to the FDA EOC.

Any FDA element alerted to an incident should promptly report the information to the FDA EOC. Upon receipt, OEM/OEO emergency coordinators or Late Duty Officer (LDO) will use contact information contained in the “Advisory Team” tab of the OEM/OEO Redbook to make the appropriate notifications.
E.2 EXTERNAL INFORMATION SHARING

Pre-identified communication pathways and information sharing methods must be established among agencies and organizations responsible for responding to a radiological incident. Effective and systematic information sharing will enhance response coordination and public messaging. FDA Centers/Offices are responsible for obtaining and updating their respective external agencies’ contact information.

E.2.1 Information Sharing Process

This section identifies specific external agencies that FDA links to for information exchange and dissemination. Not all Centers/Offices will or should be communicating with partner or external agencies. This annex specifically defines the primary FDA point of contact (POC) for each external agency or industry partner. The primary links when an IMG is activated are summarized in Figure E-2 and Table E-1.

Figure E-2 shows the primary communications links established by the IMG and its Radiological Emergency Response Unit with external agencies during a response. (An example of a full nationalized IMG organization is shown in Figure D-1.) The Radiological Emergency Response Unit consists of radiological experts who provide technical advice and serve as representatives to external agencies. Agencies external to FDA are shown in orange.

Table E-1 expands upon the communication links shown in Figure E-2 for a food and agricultural nuclear incident response and provides a summary of responsibilities for external communications. FDA roles are listed by IMG position and the supporting Center/Office. For each position, the chart lists the primary external POCs. Communication modes can include telephone, email, situation reports, inter-agency teleconferences, and 50-State teleconferences.
## Table E-1. FDA Communication Links by IMG Position and Center/Office

<table>
<thead>
<tr>
<th>IMG Position</th>
<th>Center/Office</th>
<th>External POCs</th>
<th>Notes</th>
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<td><strong>Command Staff</strong></td>
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<td>AIC</td>
<td>OEM</td>
<td>HHS SOC/ESF #8</td>
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<td>IMG PIO/FDA Joint Information Center (JIC)</td>
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<td>• Office of External Affairs (OEA)</td>
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<td>HHS Assistant Secretary for Preparedness and Response (ASPR) and Assistant Secretary for Public Affairs (ASPA)</td>
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<td>• CFSAN</td>
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<td>• Consumers</td>
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<td>• Health Professionals</td>
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<td>• National Incident Communications Conference Line (NICCL)</td>
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<td>• State Incident Communications Conference Line (SICCL)</td>
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<td>• Private Sector Incident Communications Conference Line (PICCL)</td>
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<td>• USDA (Office of the Secretary Communications and others)</td>
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<td>• CDC JIC</td>
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<td></td>
<td>• Incoming consumer calls</td>
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<td>JIC provides information out to Centers to respond</td>
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<td>LNO</td>
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<td>National Oceanic and Atmospheric Administration (NOAA)</td>
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<td>Food and Feed Safety Group</td>
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<td>• CFSAN</td>
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<td>State UC</td>
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<td>• Center for Veterinary Medicine (CVM)</td>
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<td>FRMAC</td>
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<td>• ORA Human and Animal Foods (HAF)</td>
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<td>Advisory Team</td>
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<td>• ORA Office of Communications and Quality Program Management (OCQPM)</td>
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<td>• Consumers</td>
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<td>• Health Professionals</td>
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<td>Food and Agriculture Cross Sector</td>
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<td>• Sector and consumer calls in coordination with the JIC</td>
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<td>• Also participate in Nuclear Reactors, Materials and Waste Sector call</td>
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<td>Planning Section</td>
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<td>Radiological Emergency Response Unit</td>
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<td>• OO</td>
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<td>State UC</td>
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<td>• ORA RHR</td>
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<td>• Multiple Center/Office SMEs</td>
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<td>• WEAC</td>
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<td>International Specialist</td>
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<td>HHS Office of Global Affairs</td>
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<td><strong>Logistics Section</strong></td>
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<td>Employee Health Specialist</td>
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<td>USPHS FOH Services</td>
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<td>• OCET</td>
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F. **Authorities and References**

The legal authorities that guide the structure, development, and implementation of this Nuclear/Radiological Annex include statutes and regulations, Presidential directives, national strategies, and Federal government plans and guidance. In addition, internal emergency plans and procedures augment the Nuclear/Radiological Annex and provide specific guidance to agency personnel during emergencies and disasters. In addition to the authorities and references listed in the FDA EOP, Section F, “Authorities and References,” the following authorities and references apply.\(^{73}\)

\section*{F.1 Code of Federal Regulations}


\section*{F.2 Federal Plans and Guidance}

The following list of Federal plans and guidance are applicable during a nuclear/radiological event. For a more detailed description, please see the following links:

- ESF #8 – Public Health and Medical Services  
  [https://www.fema.gov/media-library/assets/documents/25512](https://www.fema.gov/media-library/assets/documents/25512)
- ESF #10 – Oil and Hazardous Materials Response  
  [https://www.fema.gov/media-library/assets/documents/25512](https://www.fema.gov/media-library/assets/documents/25512)
- ESF #11 – Agriculture and Natural Resources  
  [https://www.fema.gov/media-library/assets/documents/25512](https://www.fema.gov/media-library/assets/documents/25512)
- Food and Agriculture Incident Annex  
- National Oil and Hazardous Substances Pollution Contingency Plan  
- NRF NRIA  
- Terrorism Incident Law Enforcement and Investigation Annex  
  [https://www.fema.gov/media-library/assets/documents/25560](https://www.fema.gov/media-library/assets/documents/25560)

\(^{73}\) This list is not exhaustive, and the associated summaries should not be used as a substitute for the authorities and references themselves.
F.3 FDA SUPPORTING DOCUMENTATION

F.3.1 WEAC Radiation Safety Manual

WEAC’s Radiation Safety Manual, Winchester Engineering and Analytical Center Policies and Procedures for Users of Radioactive Material and Radiation Emitting Devices, is issued by the WEAC Radiation Safety Committee. This document establishes procedures for the handling of radioisotopes at WEAC and other specified laboratories.

F.3.2 CFSAN Radiological Emergency Response Plan

The CFSAN Radiological Emergency Response Plan describes the responsibilities and actions to be taken by CFSAN in response to a radiological emergency. The document is Appendix W, “CFSAN Radiological Emergency Response Plan” to the CFSAN Emergency Response Plan.

F.3.3 FDA Food Protective Guidance

FDA has guidance on acceptable levels of radioactive contamination in food (called Derived Intervention Levels [DILs]) entitled, Accidental Radioactive Contamination of Human Food and Animal Feeds: Recommendations for State and Local Agencies (CDRH, 1998). This guidance provides a method for calculating the DIL for a specific radionuclide based on a limiting radiation dose to an individual consuming food contaminated at the level of a DIL for a period of one year. This guidance has been adopted by many States for their nuclear power plant safety programs and has been incorporated into EPA’s PAG Manual. FDA’s representatives on the Advisory Team, along with CFSAN radiation SMEs, assist with the interpretation of the guidance.
APPENDIX A: WINCHESTER ENGINEERING ANALYTICAL CENTER

FERN – Radiological

FDA’s WEAC radiological laboratory has analytical sample capacity during 24 hours a day, 7 days a week (24/7) operation of several hundred gamma-ray analyses per day and tens of beta- or alpha-particle analyses per week. The FERN radiological laboratories are activated by the WEAC Center Director and the FERN National Program Office when analytical surge capacity is required.

FERN will integrate the Nation’s food-testing laboratories at the local, State, and Federal levels into a network that is able to respond to emergencies involving biological, chemical, or radiological contamination of food. FERN, shared between USDA/Food Safety and Inspection Service (FSIS) and HHS/FDA, is organized to ensure ongoing Federal and State interagency participation and cooperation in the formation, development, and operation of the network (www.fernlab.org).

FERN plays a number of critical roles related to food security and food defense. These include:

- **Prevention.** FERN provides a national surveillance program that will offer early means of detecting threat agents in the American food supply.

- **Preparedness.** FERN prepares the Nation’s laboratories to be able to respond to food-related emergencies.

- **Response.** FERN offers significant surge capacity that will strengthen the Nation’s response toward widespread complex emergencies and international or inadvertent related agents in foods.

- **Recovery.** The FERN network of laboratories will enhance the ability of the country to restore confidence in the food supply following a threat or an actual emergency.

FERN includes over 165 laboratories, including 35 radiological laboratories. ORA/WEAC in Winchester, MA, is the lead FDA field laboratory for the analysis of radionuclides in foods, drugs, and medical devices.

FERN radiological laboratory contact information, laboratory capability, and capacity are maintained on the FoodSHIELD website (www.foodshield.org).

Laboratory analyses, capability, and capacity will be dependent upon the type and scope of the emergency (i.e., the source of contamination, radionuclides, foods that may be contaminated, and levels of contamination). Sample throughput is variable. For example, at the onset of the emergency, samples may be triaged through screening methods, or the use of gamma radioactivity analysis may be extremely rapid at the FDA DIL. In comparison, alpha and beta activity measurements in foods may take several days to complete.

Samples may be collected from onsite authorities or FDA, State, or local investigators and shipped to WEAC or FERN-activated laboratories.
Services Provided to FDA/ORA Personnel

WEAC is responsible for the health and safety of personnel in the field during a radiological/nuclear event. As such they provide the following services:

- **Personal Dosimetry and Monitoring**
  - Personal dosimeters are provided to field analysts and import investigators. The FDA/ORA TLD personal dosimetry program is managed through the ORA Radiation Safety Officer at WEAC.
  - WEAC provides whole-body counting (i.e., the measurement of radioactivity within the whole body). This service is offered to FDA personnel who may have ingested or inhaled gamma-emitting radioactive material. Contact the FDA/ORA Radiation Safety Officer for services.

- **Radiation Detection During Border Exams of Imported Products.** FDA/ORA investigators performing inspections at ports of entry are equipped with personal radiation detectors. SOPs (e.g., Notification and Response Procedures) are provided to the investigators. FDA/ORA/WEAC and ORA/Division of Field Investigations (DFI) manage this program.

WEAC Emergency Preparedness Plan (Extract)

WEAC promotes laboratory readiness to analyze samples for radioactive contamination and advance preparation guides through WEAC SOPs. These SOPs are readily available on the FDA/ORA/WEAC intranet website. The SOPs provide the following:

- WEAC Contact Information
- Laboratory Capability and Capacity
- Investigational Guidance
  - Sample Collection Guidance and Instructions
  - Shipping Instructions
- Radiation Safety Program Information
  - Radiation Safety Information and Guidance
  - Personal Monitoring Information and Guidance
APPENDIX B:  
ACTIVATION/NOTIFICATION OF THE ADVISORY TEAM FOR  
NUCLEAR/RADIOLOGICAL EMERGENCIES

Background
FDA is a member, and currently the chair, of the Federal Advisory Team—an interagency advisory group that can be activated during radiological incidents to assist other Federal, State, tribal, and local organizations by interpreting radiological data and making PARs to protect the public. Alert and notification activities for the Advisory Team are the responsibility of the agency chairing the team. According to the current NRF NRIA, any external organization can request the assistance of the Advisory Team by contacting the CDC EOC. Until such time as the NRIA is revised, calls to the CDC’s EOC will be transferred to the FDA EOC, who will conduct further notification of the other member agencies (i.e., USDA, EPA, and CDC) that Advisory Team assistance has been requested. Requests may also be made directly to the FDA EOC. FRMAC will also request the assistance of the Advisory Team upon its activation through the FDA EOC. This document describes the internal notification and activation procedures that the FDA EOC will follow after receiving notification from the CDC EOC or directly from an outside entity.

Scope
This document applies to any radiological incident for which Federal assistance has been requested and Advisory Team participation (deployment or consultation) is needed.

Procedure
1. As described in the 2008 version of the NRIA, the requesting call may be received by the CDC DEOC. If the incoming request is received directly by FDA from a different entity, the process begins at Step 3 below.

2. The CDC DEOC staff records the caller’s basic information (e.g., name, organization, two contact numbers, nature of request/notification, deployment logistics [if known]) and either transfers the call to the FDA After-Hours Emergency Line (1-866-300-4374 or 301-796-8240) or provides the FDA After-Hours Emergency Call Center (Call Center) number to the caller. The first method is preferred.

3. During normal duty hours (8:30 a.m. to 5:00 p.m. Eastern Time, weekdays), the incoming call is received by FDA OEO staff and referred to FDA OEM Director or OEO Director for follow-up.

4. During non-duty hours (5:00 p.m. to 8:30 a.m. Eastern Time, weekdays; 5:00 p.m. Eastern Time Friday through 8:30 a.m. Eastern Time Monday; and Federal holidays), the incoming call is received by the FDA After-Hours Emergency Call Center staff (contractor). Call Center staff record the caller’s basic information (e.g., name, organization, two contact numbers, nature of request/notification, deployment logistics [if known]).

5. The Call Center refers the call to OEO LDO, or Alternate LDO, in accordance with established Call Center procedures.

6. The Call Center sends automated email message to FDA Emergency Operations (emergency.operations@fda.hhs.gov) advising all OEO staff of the LDO notification.

7. The LDO contacts the Advisory Team Chair by telephone.

8. If the Advisory Team Chair is unavailable, the LDO contacts an alternate FDA Advisory Team member, in the order specified in the Advisory Team tab of the OEM/OEO Redbook.
9. Upon reaching the Advisory Team Chair or an alternate, the LDO confirms that they would like the other Advisory Team agencies notified, and also determines if the Advisory Team would like OEO to set up a conference call with the other Advisory Team agencies, along with the desired call time, allowing sufficient lead time for required notifications to be made.

10. The LDO notifies the OEM Director and OEO Director by telephone.

11. The OEM or OEM/OEO Director will initially make telephone contacts with the following individuals in FDA and apprise them of the events and activities as they are known:
   a. CFSAN Emergency Coordinator, with a request that they contact CFSAN’s Radiological SMEs.
   b. Emergency Response Coordinator(s), District Directors (DD) or Program Director (PD) where the incident occurred.

12. The CFSAN Emergency Coordinator will notify the EOC whether CFSAN has representation on the Advisory Team (in reachback status) and/or identify other SMEs that are available to support the Advisory Team.

13. The District Directors (DD) or Program Director (PD) will be asked to contact the RHR(s) and make arrangements for their deployment to the site of the emergency where they will coordinate with other onsite Advisory Team members.

14. The LDO sends a summary email to FDA Emergency Operations (emergency.operations@fda.hhs.gov) indicating that the call was received and successfully referred to the Advisory Team Chair or alternate member.

15. The Advisory Team representative confirms that the LDO should notify the other Advisory Team member agency EOCs (e.g., CDC, EPA, USDA) to inform them of the request for Advisory Team assistance/activation. If requested, the LDO contacts agency EOCs by email and follow-up telephone call and provides basic incident information and instructions for a conference call at the specified time. EOC telephone numbers and conference call numbers are maintained in the Advisory Team tab of the OEM/OEO Redbook.

16. The LDO notifies HHS SOC of the incoming request via email (hhs.soc@hhs.gov).

17. OEM/OEO management notifies the FRPCC Chair of the incoming request via email.

18. The LDO notifies the following individuals by email:
   a. District Directors (DD) or Program Director (PD) for the affected area.
   b. Senior Emergency Response Coordinators (SERC)
   c. All RHRs.
   d. ORA headquarters Emergency Response Coordination Group (ERCG).
   e. District Emergency Response Coordinator (DERC) for the affected District/geographic area.
   f. Director, Investigations Branch (DIB) for the affected District.
   g. CFSAN Emergency Coordinator.
   h. Counselor to the Commissioner.
   i. ORA Weekend Duty Manager (if applicable) via email.

If requested to do so by Advisory Team Chair or designated representative, the LDO includes all remaining FDA Advisory Team members and CFSAN radiological SMEs on this notification.
19. The LDO sends an information-only email to all Center/Office emergency coordinators, all SERCs, all DERCs, all DDs, and all DIBs.

20. OEM/OEO management reviews the FDA EOP, Appendix C, “FDA Mission Assignment Sub-Taskings” and initiates appropriate notifications/actions to support any contemplated FEMA/HHS mission assignment sub-taskings for Advisory Team deployment.

21. The LDO and/or OEM/OEO management maintain communications with the Advisory Team Chair or designated representative to assist with any follow-up actions required.

22. FDA EOC operational level may change according to existing procedures (see FDA EOP).74

23. For events coordinated in the FDA EOC by an IMG, the agency’s Radiological Emergency Response Unit leader will identify personnel at FDA headquarters to participate in Advisory Team conference calls and meetings to provide situational awareness and to represent the needs of the EOC for information (e.g., radiological data, maps, etc.).

NOTE: After consultation with the OEO or OEM Director, LDO duties outlined above may be distributed among other OEM managers or emergency coordinators in the interest of time.

References

- NRF
- NRF NRIA
- Advisory Team CONOPS
- FDA EOP

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74 For any radiological incident for which Federal assistance has been requested, including the Advisory Team, it is likely that the FDA EOC activation level would be increased and an IMG established.
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APPENDIX C:
PROTECTIVE ACTIONS

Protective Action Guides
PAGs are projected (estimated) dose levels to individuals in the general population at which a recommendation for protective action may be warranted. FDA has developed one set of PAGs for the ingestion pathway. The current PAGs are 5 millisievert (mSv) for committed effective dose equivalent or 50 mSv committed dose equivalent to an individual tissue or organ, whichever is more limiting. The EPA maintains PAGs for all exposure pathways, publishing them in their Manual of Protective Action Guides and Protective Actions for Nuclear Incidents (publication number 400R92001, available at EPA’s website, www.epa.gov/radiation/docs/er/400-r-92-001.pdf). EPA’s PAGs are currently in revision and a draft has been released for interim use that incorporates FDA’s current food guidance document, Accidental Radioactive Contamination of Human Food and Animal Feeds: Recommendations for State and Local Agencies, August 13, 1998 (available at www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/UCM094513.pdf).

Derived Intervention Levels
The DIL refers to the concentration of radioactive contamination present in food that, if ingested at this level over a specified time, may result in the individual receiving a projected dose equal to the PAG. FDA uses the internationally accepted units of becquerel per kilogram (Bq/kg) to express DILs, but it is common for radioactivity levels to also be expressed in units of picocuries per kilogram (pCi/kg) (1 Bq equals 27 pCi). The criteria used to calculate DILs include established dose coefficients for different age groups in the population, the fraction of food intake assumed to be contaminated, and the quantity of food consumed by that age group in the specified time period. The lowest calculated value across all age groups and target organs is designated as the DIL for that radionuclide. The assumptions used to calculate the DILs are extremely conservative. Thus, using the DILs to control food intake would realistically result in a person receiving only a fraction of the PAG dose.

FDA has calculated and published DILs for the radionuclides expected to deliver the major portion of radiation dose via the ingestion pathway during the first year following an emergency. The types of situations for which FDA has calculated DILs are: a release from a nuclear power plant, nuclear fuel reprocessing plant, or nuclear waste storage facility; the detonation of a nuclear weapon; transportation accidents; and release of material from radioisotope thermoelectric generators or heater units.

Table C-1 depicts the DILs for the nine principal radionuclides expected to be major contributors from these types of emergencies. The DIL for each radionuclide group is applied independently. The DILs apply to foods as prepared for consumption. For dried or concentrated products, such as powdered milk or concentrated juices, one should adjust by an appropriate reconstitution factor. For spices, consumed in very small quantities, a dilution factor of 10 is recommended. DILs for other types of emergencies and the other radionuclides such as terrorist activities can be determined using the same methodology.

Table C-1. Principal Radionuclides and Corresponding DILs

<table>
<thead>
<tr>
<th>Radionuclide</th>
<th>DIL (Bq/kg)</th>
<th>DIL (pCi/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sr-90</td>
<td>160</td>
<td>4,300</td>
</tr>
<tr>
<td>I-131</td>
<td>170</td>
<td>4,800</td>
</tr>
<tr>
<td>Cs-134 + Cs-137</td>
<td>1,200</td>
<td>32,000</td>
</tr>
<tr>
<td>Pu-238 + Pu-239 + Am-241</td>
<td>2</td>
<td>54</td>
</tr>
<tr>
<td>Ru-103 + Ru-106</td>
<td>Ru-103 + Ru-106 = 1</td>
<td></td>
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<tr>
<td></td>
<td>(C3 / 6,800) + (C6 / 450) &lt; 1</td>
<td>Ru-103 + Ru-106 = 1</td>
</tr>
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<td></td>
<td>(C3 / 180,000) + (C6 / 12,000) &lt; 1</td>
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Sr = Strontium      Cs = Cesium       Am = Americium   C3 and C6 = the Concentrations
I = Iodine          Pu = Plutonium    Ru = Ruthenium
Protective Action Recommendations

PARs refer to specific actions that public health officials or members of the public can take to limit the radiation dose from eating contaminated products. These recommendations focus on steps to avoid or reduce radionuclide contamination in or on food and animal feed.

Public health officials or members of the public can take protective actions before and/or after contamination is confirmed. An example of a protective action taken before confirmation of contamination is a temporary, indefinite restriction on the movement of food out of a possibly contaminated region. With confirmation of contamination and specific identification of the contaminant, other protective actions may become appropriate. Assessing the extent and impact of contamination of food includes measuring activity levels in sampled foodstuffs and comparing against the relevant DILs. Depending on the specific circumstances, individuals can take a variety of protective actions, including holding of food for a specified time period to allow contamination to diminish through normal radioactive decay, removing surface contamination by washing and diverting or destroying contaminated food. Federal agencies may receive a request to provide PARs to State and local public health authorities. If there is a request for PARs for FDA-regulated products, FDA’s RHR, the agency’s onsite liaison to the Advisory Team, works with the Advisory Team and in consultation with other FDA scientific and policy experts to develop them. PARs are given to State and local authorities via appropriate on-scene NIMS procedures. State and local officials determine specific protective actions for implementation.
A memorandum of understanding (MOU) is a critical component of any formal arrangement for cooperation between two or more entities. An MOU usually describes, in broad general terms, an area of mutual interest or concern that two or more agencies or organizations may cooperatively address. MOUs generally do not include specific information regarding detailed scope of work or the exchange of funds or human resources.

A complete listing of MOUs or other written cooperative arrangements with States, other Federal agencies and foreign government counterparts, and the World Health Organization (WHO) can be found at www.fda.gov/AboutFDA/PartnershipsCollaborations/MemorandaofUnderstandingMOUs/default.htm.

MOU with the Nuclear Regulatory Commission

On December 12, 2002, an earlier MOU between FDA and the NRC was renewed for an indefinite time. It clarifies the respective roles of each agency in regulating the safe use of radiopharmaceuticals and sealed sources or devices containing radioactive material. As a result, the NRC and FDA have established LNOs and identified key management and technical personnel for coordinating responses to emergencies or specific events of mutual interest. They have conducted joint inspections of medical events involving device failures and human- or computer-generated errors. Additionally, senior management meetings between the two agencies are conducted annually.
APPENDIX E:
FDA RESPONSIBILITIES DURING RADIOLGICAL INCIDENTS

This appendix provides an overview of the FDA components that would respond to a radiological emergency and the responsibilities each Office would have during the response and recovery phases (see Table E-1).

### Table E-1. FDA Responsibilities for Radiological Incidents

<table>
<thead>
<tr>
<th>OEM/OEO</th>
<th>Office of Foods and Veterinary Medicine (OFVM)/CFSAN</th>
<th>OFVM/CVM</th>
<th>OEA</th>
<th>ORA</th>
<th>OO</th>
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</table>
| ▪ Manage FDA’s EOC  
▪ Monitors emergency situations  
▪ Coordinates overall agency response operations | Coordinate with DHS Critical Infrastructure Food and Agriculture Sector | Serve as POC for veterinary industry | Respond to external inquiries:  
▪ Media  
▪ Consumer  
▪ Industry  
▪ Health Professionals | Represent FDA on the Advisory Team | Represent FDA on the Advisory Team |
| Serve as POC for interagency communication and coordination | Coordinate and communicate with the food and agriculture industry | Serve as POC for feed supply incidents | Coordinate with HHS ASPA, ASPR, and the CDC on public messaging | Provide RHRs |
| Communicate with the HHS SOC | Serve as POC for food and agriculture industry | Ensure safety of feed for animals (food production) | Represent FDA on NICCL, SICCL, and PICCL conference calls and other agency public affairs offices as needed | Conduct 50-State calls (OP) |
| Coordinate ESF/Recovery Support Function (RSF) MAs | Ensure the safety of food for humans | Develop agency public messaging |
| Participate in the 50-State calls | | | Serve as POC for interagency communication and coordination of messaging |
| Monitor the NICCL, SICCL, and PICCL conference calls | | | |
| Coordinate with White House/National Security Staff | | | |
| Coordinate with Advisory Team; provides scientific advice for a radiological incident | | | |
| Communicate and coordinate with WEAC and FERN | | | |

1 These functions may be performed by an IMG, when activated.
U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES
U.S. Food and Drug Administration

Pandemic Influenza Annex to the
FDA Emergency Operations Plan
Version 3.0

July 2019

OFFICE OF EMERGENCY MANAGEMENT
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A. INTRODUCTION

Pandemic influenza has the potential to cause adverse health effects for large segments of the population and animals, either directly due to exposure to the virus or indirectly as a result of the social and economic disruptions caused by the pandemic. To mitigate the consequences of pandemic influenza, the U.S. Food and Drug Administration (FDA) must be prepared to respond, with potentially limited staff, to a greater-than-normal demand and a public health need for safe and effective medical products to prevent and treat pandemic influenza, as well as other FDA-regulated products.

The Pandemic Influenza Annex to the FDA Emergency Operations Plan (EOP) establishes a single, comprehensive framework for the agency’s management of pandemic influenza emergencies. It provides the measures, operating structures, roles and responsibilities, and mechanisms for direction and coordination of FDA resources before, during, and through the recovery phase from an outbreak. The Pandemic Influenza Annex is compatible with the scalable, flexible, and adaptable Federal government emergency coordinating structures of the National Response Framework (NRF) and is consistent with the concepts, principles, and terminology of the National Incident Management System (NIMS)\(^\text{75}\) and the FDA EOP. The Pandemic Influenza Annex is to be used to assist FDA in conducting response operations for any pandemic influenza emergency.

A.1 MISSION

During an influenza pandemic emergency, FDA’s mission may expand to include performance of specific incident-related functions, such as:

- Facilitating the availability of approved and unapproved medical countermeasures, including evaluating and authorizing, as appropriate, emergency use authorizations (EUAs) for vaccines and antiviral medical countermeasures, personal protective equipment (PPE), and in-vitro diagnostics (IVDs).
- Working with manufacturers to expedite the development, licensure (if possible), and lot release of a vaccine specific to the pandemic strain.
- Monitoring the availability of critically needed FDA-regulated medical products, such as respiratory protection devices.

A.2 PURPOSE

The purpose of the Pandemic Influenza Annex is to add more detailed guidance to the FDA EOP to provide a coordinated response to the adverse impact an influenza pandemic would have on FDA and the industry and products it regulates. To accomplish this, the Pandemic Influenza Annex describes specific actions that FDA’s organizational components and personnel will take when responding to a pandemic influenza incident and enhances the agency’s emergency preparedness and response capabilities.

The Pandemic Influenza Annex:

- Is compatible with and expands on the FDA EOP.
- Defines the FDA influenza pandemic emergency operating structure and identifies the essential tasks of all FDA organizational components involved in prevention, protection, response, and recovery efforts.
- Provides mechanisms for vertical and horizontal command, control, coordination, and communication.

\(^{75}\) For more information on the NRF and NIMS, refer to the “Authorities and References” section of the FDA EOP.
• Ensures consistency with nationally recognized incident management policies and guidance.

### A.3 Scope and Applicability

This annex applies to any pandemic influenza incident for which FDA provides assistance under Emergency Support Function (ESF) #8 of the NRF, the *National Strategy for Pandemic Influenza*, the *National Strategy for Pandemic Influenza – Implementation Plan*, or under its own authority under the Federal Food, Drug, and Cosmetic Act of 1938 (FFDCA), as amended.

This annex applies to FDA’s response activities that will assist in:

1. Stopping, slowing, or otherwise limiting the spread of a pandemic to the United States.
2. Limiting the domestic spread of a pandemic and mitigating disease, suffering, and death.
3. Sustaining infrastructure and mitigating the impact on the economy and the functioning of society.

For the purpose of this annex, an influenza pandemic incident occurs when a novel strain of influenza virus emerges with the ability to infect and efficiently spread among humans. Because humans lack immunity to the new virus, a worldwide epidemic or pandemic can occur.

This *Pandemic Influenza Annex* guides FDA personnel, explaining FDA’s response activities and procedures for a cluster of novel influenza cases in verified human-to-human transmission during the World Health Organization (WHO) phases of influenza pandemic, including the alert phase and pandemic phase (see Figure A-1 on next page). WHO has developed a risk-based approach to pandemic influenza based on a continuum, which shows the phases in the context of preparedness, response, and recovery as part of an all-hazards approach to emergency risk management. In the event of an influenza pandemic emergency, FDA headquarters and field staffs will follow these response procedures.
Definitions of Phases of Pandemic Flu Within the Continuum

<table>
<thead>
<tr>
<th>Phase</th>
<th>Description</th>
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<tbody>
<tr>
<td>Interpandemic Phase</td>
<td>This is the period between influenza pandemics.</td>
</tr>
<tr>
<td>Alert Phase</td>
<td>This is the phase when influenza caused by a new subtype has been identified in humans. Increased vigilance and careful risk assessment, at local, national, and global levels, are characteristic of this phase. If the risk assessments indicate that the new virus is not developing into a pandemic strain, a de-escalation of activities toward those in the interpandemic phase may occur.</td>
</tr>
<tr>
<td>Pandemic Phase</td>
<td>This is the period of global spread of human influenza caused by a new subtype. Movement between the interpandemic, alert, and pandemic phases may occur quickly or gradually as indicated by the global risk assessment, principally based on virological, epidemiological, and clinical data.</td>
</tr>
<tr>
<td>Transition Phase</td>
<td>As the assessed global risk reduces, de-escalation of global actions may occur, and reduction in response activities or movement toward recovery actions by countries may be appropriate, according to their own risk assessments.</td>
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</table>


Figure A-1. Continuum of Pandemic Phases

A.4 PLANNING ASSUMPTIONS

The Pandemic Influenza Annex is based on the following planning assumptions:

- During the interpandemic phases, FDA will perform its routine functions.
- During the alert phase, in addition to routine functions, many FDA Offices and Centers may begin performing pandemic influenza response activities described in this annex.
- During the pandemic phase (i.e., U.S. pandemic period), routine functions may continue based on staffing availability; however, pandemic influenza response activities will take priority.
- Susceptibility to the pandemic influenza virus will be universal.
- An influenza pandemic will cause simultaneous outbreaks across the United States, limiting the ability to transfer assistance from one area to another.
- Widespread illness in communities may increase the likelihood of significant shortages of personnel (in health and non-health sectors) who provide other essential community services.

76 This continuum is according to a “global average” of cases, over time, based on continued risk assessment and consistent with the broader emergency risk management continuum.
• A pandemic will pose significant challenges to FDA employees responsible for sustaining FDA’s essential functions due to widespread absenteeism.

• An influenza pandemic may require FDA staff members to sustain some essential functions from home.

• Some information systems supporting FDA operations may not be available.

• Due to the uniqueness of their positions, not all essential employees will have a backup.

• The adverse impact of pandemic influenza on the availability of safe and effective FDA-regulated products requires an immediate response by FDA. Protection of the affected or potentially affected population is the highest priority during response operations.

• Epidemics would last 6 to 8 weeks in affected communities. Multiple waves of illness (periods during which community outbreaks occur across the country) are likely to occur with each wave lasting 2 to 3 months. As many as 40 percent of FDA employees and FDA-regulated industry employees in a geographic area could be absent from work at any point during a pandemic. This could also adversely impact the donor pool necessary for availability of blood and tissue products.

• Greater product demand and increased worker absenteeism in FDA-regulated industry could lead to shortages of commercial food products and medical products that are used to prevent and treat pandemic influenza.

• Quarantining of critical FDA-regulated facilities and large geographic areas may result. This may affect other countries, requiring extensive coordination with State, local, and foreign governments in conjunction with other Federal agencies.

• FDA U.S. Public Health Service (USPHS) Commissioned Corps personnel may be deployed, as necessary, to support U.S. Department of Health and Human Services (HHS) pandemic influenza response efforts.

• Program Directors, District Directors, and Center/Office Directors may activate their individual organizational component’s emergency operations or pandemic influenza plan without activation of the Pandemic Influenza Annex.

A.5 Activation

The Public Health and Medical Services (ESF #8) response to an outbreak of pandemic influenza will be directed by the HHS Office of the Secretary in coordination with the U.S. Department of Homeland Security (DHS), which is responsible for managing the U.S. Government (USG) response. FDA is a subordinate Operating Division (OPDIV) of HHS, and it will align its response activities with those of HHS. FDA will use the full spectrum of its resources to accomplish assigned roles, responsibilities, functions, goals, and missions.

This annex will be consulted and implemented together with the FDA EOP when there is an influenza pandemic threat abroad or in North America or a declaration of a pandemic by WHO.

A.6 Supersedence

This Pandemic Influenza Annex to the FDA EOP (2018) supersedes the FDA EOP Pandemic Influenza Annex (2014).
B. CONCEPT OF OPERATIONS
The scope of FDA emergency operations response to an influenza pandemic involves a wide array of products regulated simultaneously by the agency. The following concept of operations (CONOPS) section of this annex describes the principal authorities governing agency emergency functions that FDA conducts to perform its incident-related influenza pandemic operations.

B.1 EMERGENCY AUTHORITIES
FDA emergency authorities are listed in the FDA EOP.

B.2 EMERGENCY OPERATIONS PHASES
The following phases comprise the entire spectrum of FDA emergency operations: Prevention, Protection, Mitigation, Response, and Recovery. When a global pandemic is anticipated or does occur, FDA employees are expected to follow the activities described in the FDA EOP, Section B.2, “Emergency Operations Phases.” Although emergency operations may involve each of these phases over the course of a pandemic influenza incident, the nature and severity of an event and FDA’s organizational component(s) responding will determine the specific order, actions, and responsible parties required for each.77 The following sections describe additional resources available during FDA’s emergency operations for Prevention and Protection specific to a potential pandemic or pandemic event.

B.2.1 Prevention and Protection

B.2.1.1 Surveillance and Detection
During a pandemic influenza event, FDA may use the following surveillance and detection systems:

- Adverse Event Reporting System (AERS)
- Biological Product Deviation Reporting (BPDR) System
- Center for Devices and Radiological Health (CDRH) Product Availability (Shortages) Database
- Emergency Operations Network – Incident Management System (EON-IMS)
- Manufacturer and User Facility Device Experience (MAUDE) System
- MedWatch
- National Biosurveillance Integration System (NBIS)
- New Drug Application (NDA) Field Alert Program
- Veterinary Adverse Drug Experience (ADE) Reporting System

For a complete listing of surveillance and detection systems, refer to the FDA EOP, Table B-1.

B.2.1.2 Consumer Product Protection
In addition to the activities designated to deter intentional and unintentional acts against FDA-regulated products and to protect consumers from potential public health hazards, as described in the FDA EOP, FDA takes the following actions during an influenza pandemic:

- Expedited review of applications for vaccines, antivirals, masks, and IVDs.
- Aggressive surveillance to identify and take action to stop fraudulent promotion of products to prevent, diagnose, or treat influenza.

77 Refer to FDA EOP, Section C, “Organization and Assignment of Responsibilities,” for an overview of FDA organizational component roles and responsibilities during an emergency.
B.2.1.3 Medical Countermeasures
FDA Centers and Offices shall perform the general functions related to medical countermeasures mentioned in the FDA EOP, Section B.2.1.3, “Medical Countermeasures.”

B.2.1.4 Increased Surveillance of FDA-Regulated Pandemic Influenza Medical Products
During an influenza pandemic, FDA may increase its surveillance of FDA-regulated medical products used to prevent, diagnose, or mitigate adverse health effects of the pandemic influenza virus. Examples of such products include antivirals, vaccines, respiratory protection devices, other types of PPE, and IVD kits. Increased surveillance may include, but is not limited to, pharmacovigilance, such as tracking of vaccine adverse events or emergence of antiviral resistance; product quality, such as tracking of counterfeit products and fraudulent claims; and product availability, such as shortages of antivirals, masks, and essential medical material. The increased surveillance could be a result of conditions, such as the use of a product by a larger or different population; the use of products for other than their labeled indication; or concerns about the potential development of resistance to a product, such as antivirals or antibiotics. A specific concern is resistance developing to any antiviral drug in use for the pandemic influenza strain.

B.2.2 Response

B.2.2.1 Gain and Maintain Situational Awareness

B.2.2.1.1 Alert and Notification
FDA employees are expected to follow the activities described in the FDA EOP, Section B.2.2.1.1, “Alert and Notification.” In addition, FDA’s Office of Operations is to be notified of a pandemic event.

B.2.2.1.2 Administrative Alert and Notification
The FDA Office of Management (OM), within the Office of the Commissioner (OC), serves as the agency’s focal point for distributing information to agency staff concerning their safety and health to maintain staff agency mission support. Examples of such information include:

- Flexible workplace agreement activities.
- PPE and other medical countermeasures, such as vaccines and antivirals.
- Building closures.

B.2.2.2 Coordination of Response Actions
FDA response coordination during a pandemic will follow the FDA EOP, which will be augmented by the following coordination and response activities.

B.2.2.2.1 FDA Headquarters Response
The FDA OC Offices, Centers, and Office of Regulatory Affairs (ORA) will follow the FDA EOP, Section B.2.2.3, “Coordination of Response Actions,” which contains contacts for information dissemination, resource coordination, and priority setting. In addition, the following steps will be taken for a pandemic:

- Examine all foreign travel and restrict if necessary.
- Review plans and update all essential personnel listings.
- Ensure employees are familiar with emergency procedures.
B.2.2.2.2 Medical Countermeasures Expanded Access and EUA

FDA response regarding medical countermeasures availability and use will follow the FDA EOP, Section B.2.2.3.5, “Medical Countermeasures: EUA and Expanded Access,” and will be augmented by OC EUA standard operating procedure (SOP).

B.2.2.3 Demobilization

The FDA OC Offices, Centers, and ORA will follow the FDA EOP, Section B.2.2.4, “Demobilization,” and return to an agency-wide Task Force structure led by the Office of Counterterrorism and Emerging Threats (OCET).

B.2.3 Recovery

The FDA OC Offices, Centers, and ORA will follow the FDA EOP, Section B.2.3, “Recovery,” and return to an agency-wide Task Force structure led by OCET.
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C. ORGANIZATION AND ASSIGNMENT OF RESPONSIBILITIES

This section identifies the essential roles and functions of FDA Centers and Offices responding to a WHO alert or pandemic phase of an influenza pandemic. An essential employee is one who supports the accomplishment of a Center or Office’s pandemic influenza essential functions. Centers and Offices have provided descriptions of their pandemic influenza essential functions for inclusion in this Pandemic Influenza Annex. For an employee to be considered essential to the accomplishment of pandemic influenza essential functions, the employee need not to be assigned work solely related to pandemic influenza response.

FDA Centers and Offices work closely with OC and with industry and government partners to ensure the safety and efficacy of products for human use to prevent, diagnose, and treat the public health effects of pandemic influenza in the United States or worldwide, using novel and expeditious approaches to product regulation for optimized availability and use in all populations. During a pandemic response, issues may arise that involve more than one Center or Office. The Centers and Offices listed in this annex have additional tasks in supporting command and control operation during a pandemic that were not identified in the FDA EOP.

FDA Product Centers

FDA product Centers are responsible for the regulation of a defined set of products. Their professional staff includes both clinical and scientific experts. This expertise (analytical, laboratory, sampling procedures, subject matter expertise, and industry knowledge) is available for critical consultation should a pandemic influenza emergency occur. The National Center for Toxicological Research (NCTR) primarily conducts regulatory and applied research based on agency needs.

During an influenza pandemic, Centers are responsible for scientific evaluations and policy decisions (in cooperation with the FDA Emergency Operations Center [EOC], OCET, and ORA) in their respective program areas. FDA product Centers participate in an emergency response when the response includes, or may include, regulatory activities or products under their jurisdiction.

<table>
<thead>
<tr>
<th>Common Center Pandemic Influenza Response Activities</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alert Phase and Pandemic Phase</td>
</tr>
<tr>
<td>• Increase monitoring of the safety and effectiveness of pandemic influenza medical countermeasures.</td>
</tr>
<tr>
<td>• Manage Center responsibilities.</td>
</tr>
<tr>
<td>• Provide administrative support services (e.g., information technology [IT]/Internet/local area network [LAN]/database, telephones, safety/security, budget execution, personnel, procurement, payroll, facility management, mail).</td>
</tr>
<tr>
<td>• Post and maintain critical information on Center home page.</td>
</tr>
<tr>
<td>• Examine all domestic/foreign travel and restrict if necessary.</td>
</tr>
</tbody>
</table>

C.1 OFFICE OF THE COMMISSIONER

Several organizational components within OC will play a significant role in supporting the agency’s emergency response to a pandemic, such as OCET, and the Office of External Affairs (OEA). In addition, other OC offices, such as the Office of the Chief Counsel (OCC), Office of International Programs (OIP), and Office of Operations (OO) [Office of Financial Management (OFM) and Office of Information Management and Technology (OIMT), Office of Security, and Emergency Management (OSEM)], can provide valuable assistance during FDA’s response to a pandemic influenza emergency. A number of
agency officials will be considered essential employees specific to influenza pandemic emergency response activities. They include the following:

### OC Essential Employees

- Chief Counsel
- Chief of Staff
- Deputy Commissioner for Policy and Planning  
  - Assistant Commissioner for Planning
  - Assistant Commissioner for Policy
- Associate Commissioner for Special Medical Programs
- Associate Commissioner for External Affairs  
  - Assistant Commissioner for External Affairs
  - Assistant Commissioner for Media Affairs
- Deputy Commissioner for Foods  
  - Associate Commissioner for Food Protection
  - Chief Medical Officer and Director of Outbreaks
  - Senior Advisor to Director of Outbreaks
- Chief Scientist and Deputy Commissioner for Science and Public Health  
  - Assistant Commissioner for Counterterrorism and Emerging Threats
- Deputy Commissioner for International Programs  
  - Associate Commissioner for International Programs
- Chief Operating Officer  
  - Director, Office of Budget
  - Director, OFM
  - Director, Office of Financial Services
  - Chief Information Officer
  - Director, Office of Human Resources
  - Director, OSEM  
  - Director, Office of Security Operations
  - Director, OEM
- Associate Commissioner for Regulatory Affairs
- Director, Center for Biologics Evaluation and Research
- Director, Center for Drug Evaluation and Research
- Director, Center for Devices and Radiological Health
- Director, Center for Food Safety and Applied Nutrition
- Director, Center for Veterinary Medicine
- Director, NCTR

In addition, it is expected that each Center and Office will maintain a detailed list of essential employees, to include necessary support staff for the essential employees. The detailed lists of essential employees will be responsible for the agency’s continuity of business during an influenza pandemic and should be provided to OO.

### C.1.1 Office of the Chief Counsel

### OCC Essential Employees

- Associate General Counsel
- Deputy Chief Counsel
- Deputy Chief Counsel for Program Review
- Deputy Chief Counsel for Litigation
- Associate Deputy Chief Counsel for Drugs and Biologics
- Associate Deputy Chief Counsel for Devices, Food, and Veterinary Medicine
- Associate Deputy Chief Counsel for Litigation
  - Executive Officer
  - Senior Attorney

**One from each of these teams:**

- Counseling
- Animal Products
- Biologics
- Devices
- Drugs
- Foods
- Litigation
- Criminal
- Civil
OCC provides legal services involving the agency’s regulatory activities. FDA lawyers support the agency’s public health and consumer protection missions in two primary ways: handling litigation and providing counseling advice. OCC performs the same functions mentioned in the FDA EOP to support FDA’s response to an influenza pandemic.

C.1.2 Office of Legislation

<table>
<thead>
<tr>
<th>OL Essential Employees</th>
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</thead>
<tbody>
<tr>
<td>Assistant Commissioner for Legislation</td>
</tr>
<tr>
<td>Senior Advisor</td>
</tr>
<tr>
<td>Supervisory Congressional Affairs Specialist</td>
</tr>
<tr>
<td>Congressional Affairs Specialist (2)</td>
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</tbody>
</table>

The Office of Legislation (OL) provides Congress with information related to FDA’s pandemic influenza response activities. OL performs the same functions described in the FDA EOP to support FDA’s response to an influenza pandemic.

C.1.3 Office of Policy and Planning

The Office of Policy and Planning (OPP) is responsible for advising the Commissioner and other key agency officials on matters relating to FDA policies, rulemaking, and budgetary matters.

C.1.4 Office of External Affairs

<table>
<thead>
<tr>
<th>OEA Essential Employees</th>
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</thead>
<tbody>
<tr>
<td>Associate Commissioner for External Affairs</td>
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<tr>
<td>Deputy Director, Operations for External Affairs</td>
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<tr>
<td>Senior Management Officer</td>
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</tbody>
</table>

OEA works with outside groups in providing information related to FDA’s pandemic influenza response activities and ensures up-to-date public health advice and guidance is provided to consumers and targeted audiences.

C.1.4.1 Web and Digital Media Staff

<table>
<thead>
<tr>
<th>OMA/Web and Digital Media Staff Essential Employees</th>
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</thead>
<tbody>
<tr>
<td>Director, Web and Digital Media Staff</td>
</tr>
<tr>
<td>Senior Advisor/Public Affairs Specialist</td>
</tr>
<tr>
<td>Visual Information Specialist</td>
</tr>
<tr>
<td>Program Analysts</td>
</tr>
<tr>
<td>Technical Information Specialist</td>
</tr>
</tbody>
</table>

The OC Web and Digital Media Staff oversees the content, design, and management of FDA’s external website (www.fda.gov). OC Web and Digital Media Staff performs the same functions described in the FDA EOP to support FDA’s response to an influenza pandemic.
C.1.4.2 Office of Communications

<table>
<thead>
<tr>
<th>Office of Communications Essential Employees</th>
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</thead>
<tbody>
<tr>
<td>Assistant Commissioner for Communications</td>
</tr>
<tr>
<td>Deputy Director</td>
</tr>
<tr>
<td>Visual Information Specialist</td>
</tr>
<tr>
<td>Writer/Editors</td>
</tr>
<tr>
<td>Program Analyst</td>
</tr>
</tbody>
</table>

- Collaborate with the FDA Office of Media Affairs (OMA), HHS, Assistant Commissioner for Communications, and the Deputy Assistant Commissioner for Communications to protect consumer well-being and decrease the risk of illness, disease, or death by ensuring that the official influenza pandemic messages of the USG reach FDA stakeholders.
- Develop and distribute *Consumer Update* articles that will help stakeholders and their audiences prepare for and manage pandemic influenza issues.
- Ensure Office of Communications staff is prepared for operations, as needed, including operations from staff members’ homes and other offsite locations as necessary.

C.1.4.3 Office of Media Affairs/Media Relations

<table>
<thead>
<tr>
<th>OMA Essential Employees</th>
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</thead>
<tbody>
<tr>
<td>Assistant Commissioner for Media Affairs</td>
</tr>
<tr>
<td>Supervisory Public Affairs Specialist</td>
</tr>
<tr>
<td>Program Support Specialist</td>
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<tr>
<td>Press Officer</td>
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</tbody>
</table>

OMA serves as FDA’s primary liaison with the news media and develops much of the material FDA uses to communicate its public health and consumer protection messages to the public. OMA performs the following essential functions to support FDA’s response to an influenza pandemic:

- Collaborate with the HHS Office of the Assistant Secretary for Public Affairs (ASPA) on, and ensure appropriate integration with, the official influenza pandemic messages of the USG ([www.flu.gov](http://www.flu.gov)).
- Maintain and update emergency/after-hours SOPs and contact lists for essential media relations personnel throughout FDA.
- Ensure SOPs involving clearance of communications materials include clear indicators of when materials are ready for release and who has the authority to approve those indicators; collaborate with FDA Center communications staffs to develop boilerplate talking points that will allow quick messages to be released.
- Develop emergency backup plan that clearly indicates the primary contact, secondary contact, and contacts-on-deck in case of sickness, infrastructure failure, or other instances of unavailability.
- Ensure OIMT has assessed the technical infrastructure and is prepared for 24 hours a day, 7 days a week (24/7) operations, operations from staff members’ homes, and other offsite locations.
C.1.4.4 Office of Health and Constituent Affairs

<table>
<thead>
<tr>
<th>Office of Health and Constituent Affairs Essential Employees</th>
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</thead>
<tbody>
<tr>
<td>Director, Office of Special Health Issues</td>
</tr>
<tr>
<td>Deputy Director</td>
</tr>
<tr>
<td>Medical Advisor</td>
</tr>
<tr>
<td>Technology Information Specialist</td>
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<tr>
<td>Program Analyst</td>
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</table>

- Advise the Commissioner and other key FDA officials on matters related to patients, patient advocacy, health professionals, consumers, State and Federal activities, and industry issues.
- Assist in the planning, administration, development, and evaluation of FDA policies related to patient advocacy and health professional organizations, consumers, States, and industry on serious and life-threatening issues.
- Serve as a liaison between FDA and stakeholder organizations to educate various constituents on FDA-related issues and activities.
- Collaborate with FDA, HHS, and health professional and patient advocacy organizations to ensure the official influenza pandemic messages of the USG reach health professional and patient advocacy organizations.
- Deliver and distribute health professional articles that will help stakeholders and their audiences prepare for and manage pandemic influenza issues.
- Organize teleconferences to inform stakeholders of topics of interest for pandemic influenza (e.g., EUAs, shortages, compounding).
- Ensure staff is prepared for operations, as needed, including operations from staff members’ homes and other offsite locations, as necessary.

C.2. Office of the Chief Scientist

<table>
<thead>
<tr>
<th>OCS Essential Employees</th>
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<tbody>
<tr>
<td>Chief Scientist and Deputy Commissioner for Science and Public Health</td>
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</tbody>
</table>

The Office of the Chief Scientist (OCS) serves as the agency’s focus for scientific, medical, and related activities within OC.

C.2.1 Office of Counterterrorism and Emerging Threats

<table>
<thead>
<tr>
<th>OCET Essential Employees</th>
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<tbody>
<tr>
<td>Assistant Commissioner for Counterterrorism and Emerging Threats</td>
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<tr>
<td>Deputy Director, OCET</td>
</tr>
<tr>
<td>Director, Medical Countermeasures Initiative Regulatory Policy</td>
</tr>
<tr>
<td>Senior Advisor for Influenza</td>
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<tr>
<td>Senior Program Manager, Office Emergency Response Coordinator</td>
</tr>
</tbody>
</table>

OCET works with FDA’s medical product Centers to ensure the availability of medical countermeasures that can improve survival and mitigate and treat injuries from pandemic influenza. In addition to the functions outlined in the FDA EOP Base Plan, OCET coordinates and advises senior FDA officials on the portfolio of FDA influenza medical countermeasures through the FDA Pandemic Influenza Task Force. These include pandemic influenza vaccines, antiviral drugs, diagnostics, medical devices, and PPE such as facemasks and respirators.
**C.3  Office of Operations**

<table>
<thead>
<tr>
<th>OO Essential Employees</th>
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<tbody>
<tr>
<td>Deputy Commissioner for Operations/Chief Operating Officer</td>
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<tr>
<td>Director, Office of Equal Employment Opportunity and Diversity Management</td>
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<tr>
<td>Director, Employee Resource and Information Center</td>
</tr>
<tr>
<td>Director, Division of Technical Services</td>
</tr>
<tr>
<td>Director, Office of Business Operations and Human Capital Programs</td>
</tr>
<tr>
<td>Director, Division of Facilities Engineering and Mission Support Services</td>
</tr>
<tr>
<td>Associate Director, Division of Planning Engineering and Space Management</td>
</tr>
<tr>
<td>Associate Director, Division of Operations Management and Community Relations</td>
</tr>
<tr>
<td>Director, Office of Human Resources</td>
</tr>
<tr>
<td>Associate Director for Operations, Office of Human Resources</td>
</tr>
<tr>
<td>Associate Director for Human Capital, Office of Human Resources</td>
</tr>
<tr>
<td>Director, Client Services Division, Office of Human Resources</td>
</tr>
<tr>
<td>Director, Workforce Relations Division, Office of Human Resources</td>
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<tr>
<td>Director, OSEM</td>
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<tr>
<td>Director, OSO</td>
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<tr>
<td>Safety and Occupational Health Manager</td>
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<tr>
<td>Director, OEM</td>
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OO provides executive direction, leadership, coordination, and guidance for the day-to-day operations of FDA; manages overall budgets and resources; and oversees management and business activities agency-wide as well as OC. OO also ensures proper conduct of FDA’s administrative and financial management activities including security and emergency management.

OO will provide workforce guidance and support to FDA Offices and Centers during a pandemic influenza emergency so that the agency’s essential functions can continue with a potentially reduced number of employees. OO performs the following essential functions to support FDA’s response to an influenza pandemic:

- **Office of Business Services (OBS).** The Employee Resource and Information Center (ERIC) staff will provide critical support to FDA personnel, Offices, and Centers during a pandemic influenza emergency, including:
  - **Fleet Services:** Ensuring essential ground transportation/fleet is available and accessible to meet critical needs.
  - **Mail Services:** Ensuring critical mail pick-up and delivery.
  - Call Center support for critically needed administrative services.

- **Office of Facilities Engineering and Mission Support Services (OFEMS).** OFEMS will ensure FDA facilities are safe and available for FDA staff. Temporary facilities may be acquired if necessary. Flexibility should be built into current leases and/or custodial contracts to address cleaning and maintenance of facilities’ needs in the event of an emergency and/or have adequate supplies on hand such that FDA employees can perform these types of activities themselves, such as:
  - Building, operations, and maintenance
  - Alterations to FDA space
  - Real estate management (leasing)
### OO Response Activities

#### Alert Phase

- Interact with HHS as appropriate.
- Provide active internal planning for pandemic preparedness, including review and communication of OO’s pandemic influenza emergency response plan (ERP) to OO employees, ensuring essential personnel are equipped with necessary requirements to complete essential functions from decentralized worksites.
- Conduct disaster response training and scenario walkthroughs.
- Work with other FDA components to ensure emergency plans are ready and tested.
- Coordinate training for agency staff regarding prevention (e.g., PPE, social distancing, hygiene) and emergency operating procedures.
- Provide information to staff on personal hygiene and prevention.
- Coordinate phone numbers and services with building leasers, property management vendors, and contractors to verify their plans during the pandemic phase (i.e., backup contract services).
- Ensure essential personnel are equipped with necessary equipment to work from alternative locations, if possible.
- Advise employees planning leave or travel to take laptop computers home.

#### Pandemic Phase

- Perform all interpandemic phase and alert phase activities as appropriate and if requested by HHS.
- To the extent possible, activate and implement OO pandemic influenza Continuity of Operations (COOP) Plan as appropriate.
- Serve as agency liaison to HHS in obtaining appropriate medical countermeasures such as vaccines, antivirals, and PPE for agency staff.
- Address the health and safety of staff, in particular essential staff, that may not have the option for teleworking by providing countermeasures through occupational health services if necessary.
- Manage coordination of timekeeping functions and liaise with the Rockville Human Resources Center (RHRC) and the Program Support Center [PSC] to ensure FDA employees continue to receive compensation.
- Ensure key vendors are paid for their services to prevent termination of essential services and supplies, potentially damaging the FDA pandemic response capability.
- Ensure the FDA building closures and reopenings are updated in ERIC and posted on FDA’s intranet as circumstances warrant.
- Ensure well-being of staff, and particularly essential staff, personnel who may not have the option for teleworking.
- Ensure critical administrative functions are supported through the Call Center.
C.3.1 Office of Finance, Budget, and Acquisition

C.3.1.1 Office of Budget

In the event of pandemic influenza, FDA will likely need a supplemental budget request to support unforeseen expenses during this critical time. The Office of Budget (OB) will facilitate this process by working with senior management and Centers/Offices to identify resource needs, implementing the process to formulate the budget request, providing essential support during the review and clearance process, and responding to questions from congressional and other stakeholders.

Staffing needs will vary, based on the amount of information that OB will need to obtain and the level of analysis that is necessary. OB will likely be able to formulate a supplemental request with about three to four OB staff.

If OB is debilitated by the pandemic, it can either work with the budget analysts at HHS to assist in facilitating the supplemental budget request or use the FDA Center analysts (if available), some of whom have worked with OB in some capacity.

Because a supplemental budget request can have numerous variables and a supplemental request must be reviewed and cleared very quickly, it is difficult to provide expanded services without an additional level of investment. A supplemental budget request must be completed in its entirety.

In the event of an influenza pandemic, the budget liaison function is an essential activity to communicate budget needs and information. OB will facilitate that process by conducting outreach and coordinated communications, gathering information, and communicating with senior FDA management, Centers and Offices, HHS, Office of Management and Budget (OMB), and congressional staff.

Staffing needs will vary based on the situation; however, three to five OB analysts can facilitate the liaison process using several different channels. During an influenza pandemic, the basics of the budget liaison function can be done with OB at half the level of staffing.

Quality of work, timeframes, and milestones may slip as a result of reduced staffing. The quality and consistency control function will be important to support any budget proposals and budget information provided to others in FDA, the Administration, Congress, and stakeholders. OB will facilitate that process by using OB historical data repositories.

Staffing needs will vary based on the situation; however, the quality level of review will depend upon the number of available staff.

If OB is debilitated by an influenza pandemic, it will then work with some of the Center analysts to review budgetary documents for quality and consistency. OB can provide training to Center analysts on how it performs its quality and consistency functions, such as cross-checking budget documents and numbers.

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<tr>
<th>OB Essential Employees</th>
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<tbody>
<tr>
<td>Director</td>
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<tr>
<td>Deputy Director</td>
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<tr>
<td>Financial Advisor</td>
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<tr>
<td>Senior Analyst</td>
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<thead>
<tr>
<th>OB Response Activities</th>
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<tbody>
<tr>
<td>Alert Phase</td>
</tr>
<tr>
<td>• Track pandemic influenza budget requests and appropriations.</td>
</tr>
<tr>
<td>• Respond to questions from Congress and stakeholders.</td>
</tr>
</tbody>
</table>
• Analyze the adequacy of the FDA budget to prepare for pandemic influenza.
• Align annual budget requests to meet pandemic influenza obligations.
• If necessary, develop and advance a supplemental budget request to support unforeseen expenses related to preparing for an influenza pandemic.

**OB Response Activities**

**Pandemic Phase**

• Perform all pandemic alert activities as appropriate.

### C.3.1.2 Office of Acquisitions and Grants Services

<table>
<thead>
<tr>
<th>OAGS Essential Employees</th>
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<tbody>
<tr>
<td>Director, OAGS</td>
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<tr>
<td>Associate Director, OAGS</td>
</tr>
<tr>
<td>Director, Division of Acquisition Support and Grants</td>
</tr>
<tr>
<td>Director, Division of Acquisition Programs</td>
</tr>
<tr>
<td>Director, Division of Acquisition Operations</td>
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<tr>
<td>Director of Information Technology</td>
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The Office of Acquisitions and Grants Services (OAGS) will ensure appropriate support services and supplies are acquired to support the agency’s response to the pandemic. These include, in priority order:

1. **Purchase Cards (P-cards) (formerly International Merchant Purchase Authorization Card [IMPAC]).** OAGS manages the program and serves as the central point of contact (POC) between FDA and the central bank. Program offices use P-cards to make micropurchases (less than $3,000). In the event of an emergency, certain OAGS employees will also have P-cards with higher transaction limits.

2. **Simplified Acquisitions.** OAGS awards these through warranted contracting officers. The bulk of FDA’s acquisitions are purchase orders. There are fewer statutory requirements associated with these than with contracts, which makes them easier and faster to award than contracts. In the event that the OAGS workforce is drastically reduced or it is impossible to come into the office (e.g., under quarantine, caring for sick a family member), simplified acquisitions should be awarded whenever possible. Although the limit for a simplified acquisition is generally $100,000, this threshold may be raised to $250,000 in the event that a contingency operation is declared by the President, the Secretary of Defense, or a designee. Additionally, commercial items worth up to $5 million can be bought using simplified acquisition procedures. The absolute number of purchase orders will probably decrease or stay the same even if the simplified acquisition threshold is raised.

3. **Contracts.** OAGS contracting officers negotiate, award, and administer contracts. Whenever an acquisition exceeds the relevant simplified acquisition threshold, goods and services must be purchased using contracts. Contracts typically take longer than simplified acquisitions to award, but there are procedures, such as limiting competition on the basis of unusual and compelling urgency, that can accelerate the process.

4. **Interagency Agreements (IAGs).** OAGS negotiates and is responsible for all administrative and business-related matters associated with IAGs between FDA and other Federal agencies. In the event of severely reduced staffing, it is likely that agencies will need to leverage the resources of other offices. FDA may leverage the resources of other agencies (e.g., reaching out to other HHS OPDIVs to take over exceptionally critical FDA functions) if FDA’s staff is too small to handle
them independently. OAGS will ensure it has sufficient staff with the knowledge to execute and administer IAGs.

5. **Grants.** OAGS administers FDA’s grant program. In the event of an emergency, grants may be awarded for research or education activities that support FDA’s pandemic influenza response work.

6. **Technology Transfer.** Technology transfer encourages partnerships for collaborative research. This strengthens FDA’s research efforts and increases resources for mission-related projects. In the event of a pandemic, this is less of a priority than ensuring that mission-critical programs have the supplies they need.

### C.3.1.3 Office of Financial Operations

#### C.3.1.3.1 Office of Financial Management

<table>
<thead>
<tr>
<th>OFM Essential Employees</th>
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<tbody>
<tr>
<td>Director, OFM</td>
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<tr>
<td>Business Continuity Coordinator (Ancillary Duty)</td>
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<tr>
<td>Division Director, Financial Support Services</td>
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<tr>
<td>Division Director, Budget Execution</td>
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<tr>
<td>Division Director, Accounting</td>
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<tr>
<td>Division Director, Business Transformation, Administration and Management</td>
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### C.3.2 Office of Information Management

<table>
<thead>
<tr>
<th>OIMT Essential Employees</th>
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<tbody>
<tr>
<td>Chief Information Officer</td>
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<tr>
<td>Deputy Chief Information Officer</td>
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<tr>
<td>Director of Division of Business Partnership and Support</td>
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<tr>
<td>Deputy Director of Division of Business Partnership and Support</td>
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<tr>
<td>Director of Division of Chief Information Officer Support</td>
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<tr>
<td>Deputy Director of Division of Chief Information Officer Support</td>
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<tr>
<td>Director of Division of Infrastructure</td>
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<tr>
<td>Deputy Director of Division of Infrastructure</td>
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<tr>
<td>Director of Division of Systems</td>
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<tr>
<td>Deputy Director of Division of Systems</td>
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<tr>
<td>Director of Division of Technology</td>
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<tr>
<td>Deputy Director of Division of Technology</td>
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OIMT enables FDA’s strategic efforts to transform and improve IT systems and infrastructure needed to support critical agency operations. OIMT performs the following essential functions to support FDA’s response to an influenza pandemic:

- Implementing and enhancing common IT systems to support FDA’s response to an influenza pandemic.
- Maximizing the availability and use of IT that increases electronic access for the public, as well as the full span of FDA’s other external (e.g., public consumers, industry, government, academic) and internal customer bases, while maintaining effective security.
- Aligning IT investments to agency pandemic influenza response work.
• Consolidating, modernizing, and optimizing FDA’s IT infrastructure to strengthen its pandemic influenza response work.
• Providing the platform required for FDA to meet agency-wide IT initiatives for pandemic influenza response.

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<tr>
<th>OIMT Response Activities</th>
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<tr>
<td><strong>Alert Phase and Pandemic Phase</strong></td>
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During an influenza alert and pandemic phases, OIMT tactical activities will include:

• **Command and Control of Reporting Offices.** OIMT will define and manage the emergency while ensuring the operations of the Center and Office are maintained as required (Levels 1 to 3) and essential functions are maintained (Level 4).
• **Computer Room Facilities.** OIMT will ensure the computer room facilities function and will coordinate with PSC and building management. If cooling, generator, and power services are not available to the Parklawn Data Center, then all IT resources in the data center are also not available.
• **IT Security Services.** OIMT will ensure IT security services are available that are essential to protect FDA’s IT assets. Increased attacks on FDA’s IT resources are likely during a perceived time of weakness. Personnel will focus their efforts on security measures, such as firewall connectivity and others, ensuring that security controls for hardware, software, and telecommunications are effective and protect privacy, confidentiality, and availability of FDA data. Personnel will monitor firewall, intrusion detection, anti-malware, and security controls for critical and high-impact FDA systems, providing incident response of those potential issues that would have a severe impact on FDA operations. These services are dependent on the computer room facilities.
• **LANs and Wide Area Networks (WANs).** OIMT will ensure the availability of LANs/WANs that will be critical for both field and local users to access agency IT resources. Demand may be reduced due to fewer numbers of employees in the office. This service is dependent on the computer room facilities and IT security services. Network teams will focus their support efforts on those buildings declared operational and will not monitor or respond to network failures in buildings that are unoccupied. Networking resources can be leveraged between the Network Control Center and the LAN team.
• **Internet Services.** OIMT will maintain Internet services that are critical to the agency’s ability to communicate with regulated industry and provide access to FDA’s internal and external IT resources. The demand for these services will be greater because most staff will revert to accessing these resources remotely through the Internet. Services are dependent on outside vendors and the users’ local Internet service provider (ISP). Networking resources can be leveraged between the Network Control Center and the LAN team. If additional bandwidth is needed, the agency will request that the ISP increase bandwidth. During a pandemic event, any reengineering of the network and service segments will cease so that efforts are focused on supporting this service.
• **Active Directory.** OIMT will ensure Active Directory availability as it provides authentication services when users log on to the FDA systems. Although the demand for this service may not be great, access to IT systems will not occur if this application is down. Active Directory is dependent on computer room facilities, IT security services, LAN/WAN, and Internet services.
Telephone. OIMT will ensure telephone availability, which is a critical need. It is estimated that voice and video services will increase. Within government facilities, usage will increase substantially with the possibility of employees being ill and more working from home during a pandemic phase. This service is dependent on computer room facilities; IT security services; LAN/WAN networks; Internet services; local exchange carriers; Verizon; and the National Institutes of Health (NIH), which manage the Parklawn voice switch. This function is covered by two groups that manage Integrated System Digital Network (ISDN) and Voice over Internet Protocol (VoIP).

Email/Cell Phone Services. OIMT will ensure COOP for FDA email and smartphone services. In addition to the growing demand for email and BlackBerry services, coordination of both services is critical because each service backs the other. As of October 2009, the FDA email system is no longer operated by HHS. This change will allow OIMT to improve service and COOP for both email and smartphone services. In the event of an email failure, expert support will troubleshoot and resolve outages. In addition to communications being sent out through email and BlackBerry services, alternate communication avenues will be used, including the FDA Internet, intranet, and recorded announcements at ERIC, to ensure necessary information is communicated to employees. These alternate communication vehicles are dependent on computer room facilities (Parklawn Data Center), IT security services, LAN/WAN, Internet services, local exchange carriers, Verizon, NIH (which manages the Parklawn voice switch), and smartphone Personal Identification Number (PIN)-to-PIN services. There may be a need to have an adequate supply of additional smartphone devices to distribute to essential personnel during an influenza pandemic.

Outlook Web Access (OWA). OIMT will monitor the availability of OWA, since this is another method of accessing FDA email from non-FDA-issued computers that have access to the Internet. In order to use OWA, users will need their Remote Secure Access (RSA) SecureID token. OWA gives FDA employees greater freedom to access their email from most personally owned or public-access personal computers (PCs) because it does not require that the computer be FDA-issued or that a connection be made to the FDA network.

Help Desk Services. OIMT will manage and maintain Help Desk services where demand is expected to increase substantially as the agency executes its response plans. Although most problems may be resolved remotely, there will be fewer onsite Help Desk technicians available for support within FDA buildings. There will also be a great demand from remote logon users who have not logged into the systems in more than 6 months. An estimated 1:200 person ratio will be needed to service users. These services are dependent on building closures, numbers of users who are working in the office, and numbers who are working from home or other alternate worksites.

Secure Remote Access System (SRAS) Infrastructure. OIMT will ensure a SRAS infrastructure that will support 3,000 concurrent broadband users and about 350 dial-in users, although these numbers will increase as funding is increased. Once an order is given to shelter at home, there will be a very high demand for SRAS and, consequently, a likely surge overcapacity for dial-in service and considerable stress on broadband bandwidth. Dial-in service is not the recommended method of remote access due to the slow performance associated with this connection. This service is dependent on all essential IT infrastructure components being online, the users’ local ISP, and the telephone company having enough capacity to support the demand.

Storage Management. There are many FDA applications that will need access to data storage devices. OIMT staff may have limited capabilities to respond to all demands. A prioritized list of all applications will need to be derived to ensure adequate support of key applications.
• **Application Services.** Users will have to access FDA applications during a pandemic outbreak. OIMT staff may have limited capabilities to respond to all of the demands. A prioritized list of all applications will need to be derived to ensure adequate support resources are available. Support for this service will be dependent on the number of critical applications needed to support a response to a pandemic outbreak. Once those applications are determined, a support plan will be published that identifies the servers and storage devices that will need support. Additionally, there are other applications, although not directly supporting a pandemic outbreak, that will need to be supported, such as the Prior Notice System. These services are dependent on all critical IT resources being operational. Because the [www.fda.gov](http://www.fda.gov) website is hosted externally, OIMT will need to coordinate with the vendor to ensure continuity of service.

• **File/Print Services.** File and print services are dependent on accessibility to the network drive and will be dependent on all critical IT resources being operational. OIMT will work diligently to ensure all network drives and printers are accessible. However, there may be times, such as during a scheduled or emergency maintenance outage, that certain network drives and printers will not be available. To ensure employees have access to files and documents necessary to carry out their duties at all times, OIMT explores secure technologies, such as the IronKey device, that will allow employees to save and retrieve documents and files securely to and from an external drive.

• **Communication Services.** OIMT will continue communication services that are essential for overall information dissemination as well as for technical and project management during a pandemic influenza event. Technical IT operations information, currently issued through the OIMT News and Flash accounts, will continue to provide information on status and outages to users. OIMT can leverage the other communication officers in OO. Announcements via email are dependent on HHSMail services.

• **Contracts/Procurements Coverage.** OIMT will ensure coverage for contracts/procurements that are essential, as there may be a need to procure additional contract services or hardware/software components. This function, although important, may be performed by another OM component. This service is dependent on the availability of OAGS and key OIMT staff. OIMT can leverage other organizations to handle agency procurement needs.

• **Critical Patch Management.** Patches reduce or correct IT security threats or system problems. All efforts will be made to identify and apply patches that will address the most critical security vulnerabilities. All others will be prioritized and applied at the earliest convenience. This service is dependent on test servers and the availability of the network and server teams to load the patches.

### C.3.3 Office of Security and Emergency Management

#### C.3.3.1 Office of Emergency Management

<table>
<thead>
<tr>
<th>OEM Essential Employees</th>
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</thead>
<tbody>
<tr>
<td>Director, Office of Emergency Management</td>
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<tr>
<td>Director, Office of Emergency Operations (OEM/OEO)</td>
</tr>
<tr>
<td>Special Assistant to OEM Director</td>
</tr>
<tr>
<td>Office of Emergency Operations Staff Supervisor (2)</td>
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<tr>
<td>Emergency Coordinator (4)</td>
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</table>
OEM consists of the OEM Director (and staff) and the Office of Emergency Operations (OEO). OEM and OEO are responsible for developing crisis and emergency management policies and procedures and managing agency emergencies when they occur. OEM manages the FDA EOC.

## OEM Response Activities

### Alert Phase

- Maintain situational awareness of U.S. and worldwide pandemic influenza cases.
- Serve as agency focal point for coordination of all emergency response activities.
- Manage the agency’s EOC and activate as needed.
- Manage the EON-IMS to provide agency officials access to large volumes of data related to pandemic events.
- Create geographic information system (GIS) maps to enhance response activities.
- Manage FDA’s national Consumer Complaint Reporting system.
- Provide incident command training to OEM staff and other agency staff involved in emergency response activities.
- Participate in external emergency exercises, including pandemic influenza, and coordinate internal FDA exercises, including pandemic influenza.
- Participate in and support HHS Office of the Assistant Secretary for Preparedness and Response pandemic influenza planning activities.
- Identify agency pandemic influenza needs and goals in an agency priority setting process.
- Lead and/or assist in the development and implementation of FDA strategy for pandemic influenza.
- Lead the emergency preparedness and response component of the FDA Pandemic Influenza Preparedness Task Force.
- Lead the development of agency-wide pandemic influenza plans.
- Advise senior FDA officials on preparedness requirements for pandemic influenza.
- Provide updates to the Commissioner, other senior agency officials, and agency emergency coordinators regarding DHS, HHS, and Centers for Disease Control and Prevention (CDC) preparedness activities for potential U.S. pandemic influenza cases.
- Develop plans for FDA’s EOC and prepare for possible activation of an Incident Management Group (IMG).

### Pandemic Phase

- Perform all pandemic alert activities as appropriate.
- Give priority attention to pandemic influenza-related work and life-threatening/sustaining work associated with each of the activities listed above for pandemic alert phases.
- Communicate with other Federal, State, and local governments as they request technical and material support from FDA, and coordinate the agency’s response to the requests.
• Provide FDA liaison to HHS Secretary’s Operations Center (SOC) or other Federal EOCs as needed.
• Provide daily Situation Reports (SitReps) to the HHS SOC.
• Represent FDA on the HHS ESF #8 Pandemic Influenza Planning Group.
• Establish the FDA IMG.
• Provide leadership for Planning and Operations Sections of IMG.
• Expedite triaging of incoming consumer complaints for adverse events associated with FDA-regulated products intended for preventing, treating, mitigating, or containing pandemic influenza.
• Provide guidance on proper handling, marking, processing, and storing of classified pandemic influenza materials.
• Work with OO/OSEM/Office of Security Operations (OSO) to verify security clearances for FDA liaisons working at other government EOCs, liaisons from other agencies working in FDA’s EOC, and FDA staff working at Joint Operations Centers (JOCs) in the field.

C.4 OFFICE OF FOODS AND VETERINARY MEDICINE

Office of Foods Essential Employees

- Deputy Commissioner for Foods
- Associate Commissioner for Food Protection
- Chief Medical Officer and Director of Outbreaks
- Senior Advisor to Director of Outbreaks

C.4.1 Center for Food Safety and Applied Nutrition

CFSAN Essential Employees

- Center Director
- Deputy Directors
- Director, Office of Management Systems
- Director, Office of Food Defense Communication and Emergency Response
- Office Directors
- CFSAN COOP Leader

The Center for Food Safety and Applied Nutrition (CFSAN), in conjunction with the agency’s field staff, is responsible for promoting and protecting the public’s health by ensuring that the Nation’s food supply is safe, sanitary, wholesome, and honestly labeled and that cosmetic products are safe and labeled properly. CFSAN has the authority to regulate establishments that manufacture, process, pack, hold, or grow food involved in interstate commerce, including manufacturers, distributors, and warehouses. It also determines whether data collected by another agency or organization are adequate for FDA decisions regarding food and cosmetic issues in an emergency. CFSAN performs the following essential functions to support FDA’s response to an influenza pandemic:

• Develop and evaluate analytical methods for identifying influenza virus in foods.
• Prohibit the extra-label use of influenza antiviral drug products in animals when such use presents a risk to the public health.
CFSAN Response Activities

Alert Phase

- Develop and evaluate analytical methods for identifying influenza in food.
- Identify foods and feeds that are at elevated risk of contamination and investigate the effectiveness of food and feed processing and preparation practices for inactivating influenza viruses.
- Develop and disseminate recommendations on measures to prevent the spread of influenza virus via FDA-regulated foods.
- Activate internal planning for pandemic preparedness, including the establishment of a pandemic influenza ERP (i.e., identifying essential functions, identifying essential personnel with adequate tiers for redundancy, and equipping essential personnel with necessary requirements to complete essential functions from decentralized worksites).
- Coordinate food and feed safety activities and plans with Federal and State agencies, industry, and others.
- Food safety activities, including surveillance, inspections, and monitoring.
- Disseminate accurate information about food safety emergencies, which is vital to minimize the adverse impacts of threats or public health emergencies.

CFSAN Response Activities

Pandemic Phase

Perform all pandemic alert activities as appropriate and if requested by HHS and the Homeland Security Council (HSC):

- To the extent possible, activate and implement the CFSAN ERP as appropriate.
- Inventory human capital and determine the well-being of staff.
- Focus the Center’s remaining workforce on issues related to products associated with the ongoing situation; significant focus on ensuring food security, safety, and defense.
- Conduct outreach to the public, industry, foreign public health authorities, and other stakeholders regarding food safety.
- Provide cadre of trained CFSAN personnel for rapid deployment to the FDA EOC, to HHS, and to external organizations to provide support for contingency functions.

C.4.1.1 Coordinated Outbreak Response and Evaluation

The Coordinated Outbreak Response and Evaluation (CORE) Network is responsible for managing food-borne outbreak surveillance, response, and post-response activities related to incidents involving multiple illnesses in the United States linked to FDA-regulated human and animal food and cosmetic products. During an influenza pandemic, the FDA liaison is responsible for working with the CDC for outbreak surveillance that may involve human or animal food. CORE as a multidisciplinary team may provide subject matter expertise, as needed, to the Centers. CORE would not coordinate or manage FDA’s
response. If the agency activates the FDA EOC, CORE may provide resources, to the extent possible, to support the agency’s incident management response.

**C.4.2 Center for Veterinary Medicine**

The Center for Veterinary Medicine (CVM) ensures the safety, efficacy, and quality of drugs for animals, including food-producing and companion animals, animal food and feed, and medical devices used on animals potentially threatened by an influenza outbreak. CVM performs the following essential functions to support FDA’s response to an influenza pandemic:

- Maintaining essential international program activities.
- Collaborating with public health agencies (e.g., CDC, HHS, and the U.S. Department of Agriculture [USDA]) regarding feed contaminant, tissue residue programs, and other monitoring programs for meat and poultry involved in influenza outbreaks.
- Providing advice in the assessment of animal drug or feed product possibly affected by influenza.
- Providing advice to pet owners regarding animal safety measures related to influenza.

**CVM Response Activities**

<table>
<thead>
<tr>
<th>Alert Phase</th>
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<tbody>
<tr>
<td>- Coordinate food and feed safety activities and plans with Federal and State agencies, industry, and others.</td>
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<tr>
<td>- Identify foods and animal feeds that are at elevated risk of contamination and investigate the effectiveness of food and feed processing and preparation practices for inactivating influenza viruses.</td>
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<tr>
<td>- Prohibit the extra-label use of influenza antiviral drug products in animals when such use presents a risk to the public health.</td>
</tr>
<tr>
<td>- Manage CVM laboratory capabilities and provide this information to the FDA EOC.</td>
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<tr>
<td>- Develop and disseminate recommendations on measures to prevent the spread of influenza virus via FDA-regulated foods and animal feeds.</td>
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<thead>
<tr>
<th>Pandemic Phase</th>
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<tbody>
<tr>
<td>- Perform all pandemic alert activities as appropriate.</td>
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C.5 Office of Medical Products and Tobacco

C.5.1 Office of Special Medical Programs

<table>
<thead>
<tr>
<th>OSMP Essential Employees</th>
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<tr>
<td>Associate Commissioner Special Medical Programs</td>
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C.5.1.1 Office of Combination Products

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<thead>
<tr>
<th>OCP Essential Employees</th>
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<tbody>
<tr>
<td>Director</td>
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<tr>
<td>General Attorney (2)</td>
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<tr>
<td>Biologist (2)</td>
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The Office of Combination Products (OCP) ensures the prompt assignment of combination products to FDA Centers, the timely and effective premarket review of such products, and consistent and appropriate post-market regulation of like products subject to the same statutory requirements to the extent permitted by law. OCP performs the same functions mentioned in the FDA EOP to support FDA’s response to an influenza pandemic.

C.5.2 Center for Drug Evaluation and Research

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<tr>
<th>CDER Essential Employees</th>
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<tbody>
<tr>
<td>Center Director</td>
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<tr>
<td>Deputy Director</td>
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<tr>
<td>Director, Office of Regulatory Policy</td>
</tr>
<tr>
<td>Director, Office of Communications</td>
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<tr>
<td>Director, Office of Strategic Programs</td>
</tr>
<tr>
<td>Director, Counterterrorism and Emergency Coordination Staff</td>
</tr>
<tr>
<td>Director, Office of Compliance</td>
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<tr>
<td>Director, Office of Pharmaceutical Quality</td>
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<tr>
<td>Director, Office of Surveillance and Epidemiology</td>
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<tr>
<td>Director, Office of Translational Sciences</td>
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<tr>
<td>Director, Office of New Drugs</td>
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<tr>
<td>Deputy Director, Office of New Drugs</td>
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<tr>
<td>Director, Office of Antimicrobial Products</td>
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<tr>
<td>Associate Director, Drug Shortage Staff</td>
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<tr>
<td>Director, Office of Management</td>
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</table>

The Center for Drug Evaluation and Research (CDER) ensures the safety, efficacy, and quality of drugs and therapeutic biologic agents for use as medical countermeasures in response to outbreaks of an influenza virus. CDER performs the following essential functions to support FDA’s response to an influenza pandemic.

**CDER Response Activities**

<table>
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<th>Alert Phase</th>
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- Perform drug approval activities (i.e., review of investigational and new drug applications, associated labeling and advertising, and guidance for field inspections).
• Perform drug safety activities, including adverse event monitoring, drug product quality activities, and post-market surveillance.

• Perform compliance activities, including regulatory surveillance assessments and actions and coordination of actions involving recalls, shortages, tampering, fraudulent or counterfeit products, and inspections.

• Perform emergency response coordination/activities for all CDER-regulated products, liaison to the Strategic National Stockpile (SNS), maintenance of the CDER Situation Room (CSR), program responsibility for classified documents and security clearances, and coordination of EUAs.

• Monitor for drug shortages.

• Facilitate communication with manufacturers of pandemic influenza products on issues of availability, adverse event reporting, and potential new products.

• Collaborate with HHS regarding stockpile issues, including labeling, appropriate usage, shelf-life and expired product considerations, product performance, pharmacovigilance and monitoring, use in special populations, and other evolving issues.

• Process pre-EUAs in anticipation of declared emergencies.

• Activate internal planning for pandemic preparedness, including the establishment of a Continuity of Operations Plan (COOP); identifying essential functions and essential personnel with adequate tiers for redundancy; and equipping essential personnel with necessary equipment/procedures to complete essential functions from decentralized worksites.

### CDER Response Activities

#### Pandemic Phase

• Continue to perform drug regulatory review activities (i.e., review of investigational and new drug applications, associated labeling and advertising, and guidance for field inspections).

• Perform all interpandemic phase and pandemic alert activities as appropriate and if requested by HHS and the HSC.

• To the extent possible, activate and implement the CDER COOP, CDER Pandemic Administrative Operating Plan, and other ERPs as appropriate.

• Inventory human capital and the capability of staff to work.

• Focus the Center’s remaining workforce on issues related to products associated with the ongoing situation, significant life-saving products, and new generic drugs to prevent shortages and respond to significant adverse event incidents, EUA requests, and emergency response coordination issues.

• Conduct public outreach to industry, foreign public health authorities, the public, and other stakeholders regarding availability and regulatory status of antiviral drugs.
C.5.3 Center for Biologics Evaluation and Research

CBER Essential Employees

Center Director
Deputy Director
Associate Directors
Emergency Operations Coordinator
International Affairs Advisor
Program Manager for Pandemic Influenza
Senior Advisor for Counterterrorism/Medical Countermeasures
Director, OM
Director, Office of Vaccine Research and Review
Director, Office of Blood Research and Review
Director, Office of Compliance and Biologics Quality
Director, Office of Communications, Outreach, and Development
Director, Office of Biostatistics and Epidemiology
Director, Office of Cellular, Tissue, and Gene Therapies
CBER Office of Information Management Liaison

The Center for Biologics Evaluation and Research (CBER) ensures the safety, efficacy, and quality of biological products potentially used as medical countermeasures during a pandemic influenza emergency. CBER performs the following essential functions to support FDA’s response to an influenza pandemic.

CBER Response Activities

Alert Phase

- Participate, through the WHO Global Influenza Surveillance and Response System (GISRS) to obtain, identify, and characterize emergent influenza viruses with pandemic potential.
- Work with WHO Collaborating Centres (CCs) and WHO Essential Regulatory Laboratories (ERLs) to generate, calibrate, and cross-calibrate reference reagents for vaccine formulation as necessary.
- Continue internal capacity building efforts, including development of novel reassortants (animal H1, H2, H5, H7, H9, etc.) suitable for vaccine production, methods development to improve assays for clinical assessment, and assays necessary for product release.
- Provide guidance to sponsors regarding new and/or advanced product development and clinical trials research, including novel vaccines and therapies.
- Conduct post-marketing surveillance and enforcement activities relating to biological products in accordance with pandemic guidance as appropriate.
- Industry outreach for purposes of increasing vaccine production and fill/finish capacity to meet pandemic demand.
- Provide CBER management that will carefully examine all domestic and foreign travel to restrict if necessary.
- Collaborate with HHS, other parts of the USG, and other international organizations, such as WHO, on the framework and method for appropriate selection of pandemic reference strains for vaccine production.
• Collaborate with HHS regarding stockpile issues, including labeling, appropriate usage, shelf-life considerations, product performance, pharmacovigilance and monitoring, use in special populations, and other evolving issues.

• Provide active internal planning for pandemic preparedness, including the establishment of a pandemic influenza ERP and continual updates as needed, identifying essential functions and essential personnel with adequate tiers for redundancy, and equipping essential personnel with necessary requirements to complete essential functions from decentralized worksites.

• To the extent possible, assess the availability of biological products and take steps to divert or mitigate the impact of shortages.

• Provide training to staff regarding prevention (e.g., PPE, social distancing, hygiene) and emergency operating procedures.

• Play an integral part in any global harmonization of regulatory processes within FDA’s existing framework for pandemic activities, should this be necessary.

### CBER Response Activities

#### Pandemic Phase

• Engage in pandemic activities as appropriate and if requested by HHS and the HSC.

• To the extent possible, activate and implement the CBER Pandemic Influenza COOP Plan and other ERPs as appropriate.

• To the extent possible, ensure well-being of staff and, particularly, essential staff personnel who may not have the option for teleworking (such as those involved in laboratory work supporting vaccine production, quality control, and release) by the provision of countermeasures through occupational health services in coordination with OC’s OM.

• Respond to and process EUA requests after a declaration of public health emergency by the HHS Office of the Secretary.

• Using a risk-based approach, prioritize allocation of review and inspectional resources for licensure and emergency use of medical countermeasures, as well as other essential activities that will confer the maximum and most beneficial public health impact.

• Ensure timely information flow within CBER, as well as other organizations within the USG.

• Facilitate dialogue and information exchange with industry and international entities, as well as participation in global efforts to develop consensus approaches, to the extent possible, for clinical trial designs and adverse event reporting that can guide vaccination strategies.

• Conduct public outreach to domestic and foreign public health authorities, the public, and other stakeholders regarding pandemic vaccines and other biological products.
C.5.4 Center for Devices and Radiological Health

CDRH ensures the safety, efficacy, and quality of medical devices used during a pandemic influenza emergency. CDRH performs the following essential functions to support FDA’s response to an influenza pandemic.

- Educate device users on how to use influenza-related devices safely and effectively.
- Communicate influenza-related device activities and information to industry and consumers.
- Process pre-EUAs in anticipation of declared emergencies.
- Facilitate device development, regulatory review, and production of devices that may be needed during an influenza pandemic.
- Conduct post-market surveillance to monitor the safety and effectiveness of influenza-related devices (including devices used for diagnosis, devices used to administer therapeutics, PPE, and devices used for supportive care).
- Lend technical expertise and assistance to international efforts to prepare for a potential influenza pandemic.
- Support efforts to ensure an adequate supply of influenza-related devices through cooperative interactions with manufacturers and distributors and coordination with the SNS to determine the adequacy of stocks and actions required to meet targeted amounts.
- Use an emergency shortages database to identify and monitor needed supplies.
- Collaborate with public health agencies regarding product stockpile issues, appropriate usage, labeling, product performance, monitoring, use in special populations, and other evolving issues.
- Collaborate with HHS regarding stockpile issues related to medical devices.
- Activate internal plans for pandemic preparedness in regard to staff communication and readiness, including teleworking during the pandemic.
- Provide training for staff on prevention and protection techniques for the influenza pandemic that must be followed in the workplace.
C.6 Office of Global Regulatory Operations and Policy

C.6.1 Office of International Programs

<table>
<thead>
<tr>
<th>OIP Essential Employees</th>
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<tbody>
<tr>
<td>Deputy Commissioner for International Programs</td>
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<tr>
<td>Associate Commissioner for International Programs</td>
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<tr>
<td>Deputy Director, OIP</td>
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</table>

OIP is responsible for coordinating and, if appropriate and permitted, communicating appropriate emergency-related information to foreign governments and international organizations, such as WHO. OIP also receives information and requests for information (RFIs) from foreign governments. OIP performs the following essential functions to support FDA’s response to an influenza pandemic:

- Maintain contacts and efficient channels for communications with FDA counterpart agencies and FDA staff in other countries to assess the impact of the pandemic on FDA and advise on appropriate actions.
- Monitor and report on relevant international development and resources.

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<th>OIP Response Activities</th>
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<tr>
<td>Alert Phase</td>
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- Exchange information with foreign counterparts to ensure compliance with FDA laws, regulations, and policies and consistency in responses to an influenza pandemic.
- Monitor and report on relevant international developments and resources.
- Coordinate and advise on international technical cooperation, assistance, and capacity-building activities.
- Establish policies and procedures pertaining to international travel and processing international travel requests.

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<th>OIP Response Activities</th>
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<td>Pandemic Phase</td>
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- Perform all pandemic alert activities as appropriate.
- Process EUAs for products after a declaration of public health emergency by the HHS Office of the Secretary.
- Conduct post-market surveillance to monitor the safety and effectiveness of influenza-related devices (including devices used for diagnosis, devices used to administer therapeutics, PPE, and devices used for supportive care).
- Support efforts to ensure an adequate supply of influenza-related devices through cooperative interactions with manufacturers and distributors and coordination with the SNS to determine the adequacy of stocks and actions required to meet targeted amounts.
- Use an emergency shortages database to identify and monitor supplies of certain devices that have the potential to be in demand, but in short supply during influenza outbreaks.
• Conduct international pandemic alert activities.
• Follow the Pandemic Influenza Annex to the FDA EOP and other relevant advice.
• If staff is limited, OIP functions will be restricted to:
  - Communicating with foreign governments and international organizations, including foreign embassies and consulates in the United States.
  - Exchanging information with foreign counterparts to ensure compliance with FDA laws, regulations, and policies and consistency in responses.
  - Implementing policies and procedures pertaining to international travel and processing international travel requests; OIP will process only international travel that is consistent with requirements and limitations pertaining to the pandemic situation.

C.6.2 Office of Regulatory Affairs

<table>
<thead>
<tr>
<th>ORA Essential Employees</th>
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<tbody>
<tr>
<td>Associate Commissioner for Regulatory Affairs</td>
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<tr>
<td>Deputy Associate Commissioner for Regulatory Affairs</td>
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<tr>
<td>Associate Commissioner for Medical Products and Tobacco</td>
</tr>
<tr>
<td>Associate Commissioner for Human and Animal Foods</td>
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<tr>
<td>Program Directors</td>
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<td>Program Senior Advisor</td>
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<td>Office Director</td>
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<td>District Director</td>
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<td>Program Division Director</td>
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<td>Laboratory Director</td>
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<tr>
<td>Program Director of Investigations Branch</td>
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<tr>
<td>Program Director of Compliance Branch</td>
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<tr>
<td>Program Compliance Officer</td>
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<td>Import Compliance Officer</td>
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<tr>
<td>Legal Instrument Examiner</td>
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<tr>
<td>Administrative Officer</td>
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<tr>
<td>Supervisory Consumer Safety Officer</td>
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<tr>
<td>Emergency Response Coordinator</td>
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<tr>
<td>Laboratory Supervisor</td>
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<tr>
<td>Special Assistant to District Director</td>
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<tr>
<td>Commissioned Corps Officers</td>
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<tr>
<td>Special Agent in Charge (SAIC)</td>
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<tr>
<td>Office of Criminal Investigations (OCI) Headquarters/Investigative Operations Division</td>
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<tr>
<td>SAIC OCI Headquarters/Administrative Operations Division</td>
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<tr>
<td>SAIC OCI Office of Internal Affairs</td>
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<tr>
<td>Senior Operations Manager, OCI/Headquarters</td>
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<tr>
<td>COOP Coordinator</td>
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<tr>
<td>Director of Import Operations Branch</td>
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<tr>
<td>Investigator/Consumer Safety Officer</td>
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<tr>
<td>Criminal Investigator/Special Agent</td>
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<td>Consumer Complaint Coordinator</td>
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<td>Program Recall Coordinator</td>
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<tr>
<td>Director of Cooperative Programs</td>
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<tr>
<td>Specialists/Cooperative Programs (Milk, Shellfish, Retail, Radiological Health)</td>
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<tr>
<td>Laboratory Analyst</td>
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<tr>
<td>Laboratory Branch Director</td>
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<tr>
<td>Associate Director for Business and Safety Operations</td>
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<tr>
<td>Supervisory Industrial Hygienist</td>
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<tr>
<td>Industrial Hygienist</td>
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<tr>
<td>Staff Assistants</td>
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<tr>
<td>National Health Fraud Coordinator</td>
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<tr>
<td>Public Information Branch Representative/Manager</td>
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<tr>
<td>Director, Headquarters/OCI Public Affairs Specialist</td>
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<tr>
<td>Program Support (as needed for response)</td>
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<tr>
<td>Timekeeper</td>
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<td>Sample Custodian</td>
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</table>

ORA, led by the Associate Commissioner for Regulatory Affairs (ACRA), protects consumers and enhances public health by maximizing compliance of FDA-regulated products and minimizing risk associated with those products. ORA is the lead office for all agency field activities. ORA maintains a database of FDA-regulated establishments that enables FDA to rapidly identify establishments adversely affected by a pandemic influenza incident. In the event of product shortages, ORA’s establishment
database can assist the agency in identifying firms with needed production capabilities or product inventory. ORA performs the following essential functions to support FDA’s response to an influenza pandemic:

- Investigating and pursuing enforcement action against fraudulent or counterfeit pandemic influenza products.
- Initiating surveillance and follow-up activities for fraudulent pandemic products sold over the Internet.
- Conducting post-market surveillance activities related to vaccines (team biologics).

### ORA Response Activities

#### Alert Phase

- Conduct domestic and foreign inspections; import entry review, field exams, sample collections, and analyses; and take appropriate enforcement actions against violative firms or products of heightened concern because of their relation to pandemic influenza (for instance, vaccines and influenza drugs) to ensure compliance with applicable regulations.
- Investigate and pursue enforcement action against fraudulent or counterfeit pandemic influenza products, both domestic and foreign.
- Provide incident command training to essential staff and other staff involved in emergency response activities.
- Participate in internal FDA and external pandemic influenza emergency exercises, including pandemic influenza.
- Ensure staff members have equipment and supplies needed in order to work from home.
- Participate in educational outreach activities for industry, FDA regulatory counterparts, and other external customers.
- ORA management to carefully examine all foreign travel and restrict if necessary.
- Maintain situational awareness of pandemic influenza cases worldwide through OEM.
- Activate internal ORA planning for pandemic preparedness, including the establishment of a pandemic influenza ERP; identify essential functions and essential personnel with adequate tiers for redundancy; and equip essential personnel with necessary equipment to complete essential functions for decentralized worksites.

#### Pandemic Phase

- All activities stated above also apply during an alert phase, as appropriate, and if requested by HHS and the HSC.
- Intensify efforts to identify fraudulent, harmful, or ineffective products that are promoted for preventing, diagnosing, or treating pandemic influenza.
- Give priority attention to pandemic influenza-related work and life-threatening/sustaining work associated with emergency response coordination activities.
- To the extent possible, activate and implement the pandemic influenza ERP, COOP Plan, and other ERPs as appropriate.
C.6.2.1 Office of Regulatory Affairs Laboratories

ORA laboratories will provide analytical support for agency efforts to identify counterfeit or fraudulent pandemic influenza products and to support their removal from the marketplace. ORA laboratories will also participate in shelf-life extension and related programs associated with the maintenance of national stockpiles of critical influenza products.
D. DIRECTION, CONTROL, AND COORDINATION

This section provides a description of Incident Command System (ICS) positions and groups, which are specific to a pandemic influenza response and describes their primary roles and responsibilities.

FDA response operations for major incidents, such as pandemic influenza, are based on the concepts, practices, and principles of NIMS and its organizational structure, the ICS. Among other things, NIMS prescribes what types of information FDA should provide the HHS SOC and in what format, including SitReps and Incident Action Plans (IAPs). NIMS also requires the use of universal terminology to ensure a common understanding of requirements and requests in an interagency environment. The ICS organizational structure provides standard nomenclatures for both teams and individual positions. For a complete listing of the FDA ICS structure (IMG)\textsuperscript{78} refer to FDA EOP, Section D, “Direction, Control, and Coordination.”

Expansions of the FDA response structure for a pandemic influenza response may vary according to numerous considerations and operational factors. ICS offers flexibility in determining the right structural approach for the specific circumstances of the incident at hand. Dependent on the incident objectives established and strategies developed, the IMG may include Logistics and Finance/Administrative as part of the response and expand FDA’s command structure Operations Section, including any one of the groups described in Section D below (see Figure D-1).

![Figure D-1. Possible Pandemic Influenza IMG Structure](image)

\textsuperscript{78} For more information on specific roles and responsibilities of each ICS position, refer to the FDA Incident Management Handbook.
D.1 COMMAND STAFF
The FDA Command Staff includes a Public Information Officer (PIO), Liaison Officers (LNOs) and Safety Officer (SO). Technical Advisors may be assigned to the Planning Section or to the Command Staff working directly for the Agency Incident Coordinator (AIC). Technical Advisors can include the Science/Medical Advisor, EUA/Product Interface Advisor, International Advisor, and Legal Advisor.79

D.1.1 Pandemic Influenza-Specific Liaisons
When activated, the FDA EOC serves as the agency-wide focal point for pandemic influenza emergency operations, coordination, and communications. External liaisons from the CDC or other agencies may be present at the FDA EOC. FDA may also send liaisons to the coordinating agency for the event or participate in the response coordination if an incident or unified command is set up.

D.1.2 Legal Advisor

D.1.2.1 Legal Group
The Legal Group works to address the following issues related to pandemic influenza in support of the Legal Advisor:

- Adverse event reporting guidance.
- Fraudulent products warning letters and enforcement actions.
- FDA-regulated products used during an influenza pandemic (i.e., export and import, overriding patents, and public informational statements).
- EUAs.
- Occupational issues (e.g., agency recommendations for employees, ensuring harmonization of multiple guidance for employees provided by multiple entities).
- Preparation for congressional hearings.

D.2 GENERAL STAFF
The General Staff is responsible for the functional aspects of the FDA ICS. The General Staff consists of the Operations, Planning, Logistics, and Finance/Administration Sections, which comprise designated FDA headquarters and field emergency staffs. Individual position assignments within these Sections are discussed in the FDA EOP.

D.2.1 Operations Section
The Operations Section is responsible for coordinating all agency activities related to regulated products in response to an influenza pandemic event. Figure D-1 depicts FDA’s Operations Section during a pandemic influenza response. The groups listed below could possibly be established when the number of resources exceeds the manageable span of control of the IMG and the Operations Section Chief. Groups are established to divide the incident into functional areas of operation. Additional levels of supervision may also exist below the Group level.

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79 See the FDA EOP, Section D, for a description of the Technical Advisor positions.
D.2.1.1 Drug Group
The Drug Group addresses drug issues, such as drug product availability, adverse event reporting, labeling changes, whether EUAs for drugs are appropriate, and what restrictions should be placed on uses authorized under EUAs related to an influenza pandemic operational response. The following are some potential teams, led by CDER staff, that may be needed to manage mitigation and recovery issues.

D.2.1.1.1 Antiviral Team
The Antiviral Team addresses issues concerning the availability of antiviral drugs and EUA requests and manages adverse event reporting and monitoring.

D.2.1.1.2 Expired Drugs Team
The Expired Drugs Team deals with such issues as use of expired antiviral drugs and drugs out of specification, including those in the SNS. This team would have input in discussing whether an EUA should/can include private sector stockpiles (e.g., hospital associations, large employers) or whether a separate authorization is needed. The Expired Drugs Team would also monitor labeling and Shelf Life Extension Program (SLEP) issues related to the event.

D.2.1.1.3 Drug Shortage Team
The Drug Shortage Team is responsible for reporting on the availability and surge capacity for antiviral drugs, monitoring commercial inventory, and working with manufacturers of influenza drug products to prepare for possible increases in demand. If needed, the team would contact manufacturers of intravenous antibiotics, intravenous fluids, and other critical care drugs to prepare for possible increased needs.

D.2.1.1.4 Adverse Drug Events Team
The Adverse Drug Events Team assists with the monitoring of adverse drug events specifically to identify undesired side effects and unexpected adverse effects from a drug product used to treat influenza.

D.2.1.2 Biologics Group
The Biologics Group addresses vaccines, blood, and other biologics issues as part of an influenza pandemic operational response. Some potential teams, with team leaders from CBER staff, may be needed to manage mitigation and recovery issues.

D.2.1.2.1 Vaccine Team
The Vaccine Team works with U.S.-licensed manufacturers of influenza vaccines concerning the regulatory and scientific aspects of the development and licensure of vaccines for the novel influenza virus. They participate in various recurring meetings facilitated by the Biomedical Advanced Research and Development Authority (BARDA), National Vaccine Program Office (NVPO), CDC, WHO, and international regulators, such as the European Medicines Agency (EMEA) and Health Canada.

In addition, the Vaccine Team works to address any issues related to:

- Preparation of reference strains from which manufacturers can make viral seeds for vaccine production.
- Production of reagents to test vaccine potency.
- Providing strains for assays and manufacture as necessary.
- Standardization of clinical trials.
- Vaccine production.
• Discussion with manufacturers, when needed, regarding the information needed to potentially support the use of unapproved vaccines in a declared emergency under an EUA.

**D.2.1.2.2 Blood Team**

The Blood Team is responsible for providing guidance related to blood and tissue donations and use of such products when impacted by a pandemic. The team also works with outside groups to address any blood shortages that might arise.

**D.2.1.3 Medical Device Group**

The Medical Device Group would manage issues related to an influenza pandemic operational response. Potential units, with team leaders from CDRH staff members, that may be needed to manage mitigation and recovery issues are the IVD Team, PPE and Other Devices Team, and Shortage Team.

**D.2.1.3.1 In-Vitro Diagnostics Team**

The IVD Team performs the following:

• Respond to triage EUA requests.
• Respond to inquiries regarding EUA requests from:
  – The CDC, U.S. military, and domestic and international commercial sources for IVDs claiming to subtype or detect the novel viral strain or to detect virus mutant variants.
  – Manufacturers regarding instruments to be used in the development of IVDs.
• Clear tests and instruments for the availability of detection of the novel influenza, including rapid tests and other influenza testing-related products, such as viral transport media/swabs.
• Monitor reagent and system shortages on a short-term (daily or weekly) basis as well as a long-term (monthly) basis.
• Monitor claims for detection and differentiation of the influenza virus.
• Review data and labeling of cleared or approved devices for which a company proposes new claims and changes.
• Work with the CDC to monitor availability of new influenza subtype panels from CDC/American Type Culture Collection (ATCC) for use by assay developers.
• Conduct rapid review of any promising influenza devices capable of differentiating novel subtypes, particularly those that are rapid or point-of-care tests, and those for detection of mutant variants of the virus.
• Address fraudulent claims by monitoring websites and complaints regarding claims to diagnose novel influenza and issuing warning letters as appropriate.
• Support the development of laboratory-developed tests for virus diagnosis.

**D.2.1.3.2 Personal Protective Equipment and Other Devices Team**

The PPE and Other Devices Team performs the following:

• Exchange information with CDC/SNS, States, and manufacturing sector and provide real-time monitoring of consumption/utilization rates of PPE during emergencies.
• Respond to State and private sector inquiries about perceived shortages during public health emergencies.
• Anticipate and plan for future surges in demand for N95 devices.
• Track and trace status of N95 devices released from SNS.
• Meet with manufactures about the development of new PPEs.
• Address shortages of PPE by working with industry to update current manufacturing capabilities and assist with long-term planning by the manufacturing sector to meet demands for a potential surge.

D.2.1.3.3  Shortage Team
The Shortage Team would be created if shortages of IVDs, PPEs, and other devices are too extensive to be managed by the above units alone.

D.2.2  Logistics Section

D.2.2.1  Employee Health Advisor
The Employee Health Advisor is a staff member of the OO Employee Safety and Environmental Management Staff (ESEMS) who addresses the need for appropriate health protection for high-risk FDA employees. The Employee Health Advisor works with HHS and the USPHS Federal Occupational Health (FOH) Services to ensure adequate medical countermeasures are made available to protect FDA responders.
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E. COMMUNICATIONS AND INFORMATION MANAGEMENT

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F. AUTHORITIES AND REFERENCES

F.1 NATIONAL STRATEGIES

National Strategy for Pandemic Influenza

On November 1, 2005, President George W. Bush issued a National Strategy for Pandemic Influenza that called for comprehensive and coordinated pandemic preparedness planning at all levels of government and the private sector. It outlines how the Nation intends to prepare, detect, and respond to a pandemic. This strategy describes the important roles to be played not only by the Federal government, but also by State and local governments, private industry, international partners, and individual citizens. HHS was identified as the lead department for medical response.


National Strategy for Pandemic Influenza – Implementation Plan

In May 2006, the HSC published the National Strategy for Pandemic Influenza – Implementation Plan. The introductory letter signed by President George W. Bush cited more than 300 critical actions to address the threat of pandemic influenza. Chapter 6, Protecting Human Health, includes actions on (1) achieving national goals for producing and stockpiling vaccine and antiviral medications; and (2) prioritizing and distributing limited supplies of vaccine and antiviral medications. It details how the Nation is currently preparing for, and how it detects and responds to, a potential pandemic influenza outbreak. www.flu.gov/professional/federal/pandemic-influenza-implementation.pdf

F.2 FEDERAL PLANS AND GUIDANCE

HHS Pandemic Influenza Plan, Part 1 – Strategic Plan

The HHS Pandemic Influenza Plan, Part I – Strategic Plan (November 2005) contains a summary of major pandemic response roles for HHS officials, agencies, and divisions.


The major pandemic response roles for FDA include the following:

- Regulating manufacturing processes.
- Evaluating and licensing pandemic vaccines.
- Evaluating and approving antiviral drugs for influenza.
- Facilitating the development, evaluation, and clearance or approval of diagnostic tests and devices.
- Preparing reagents to standardize the potency of inactivated influenza vaccines.
- Preparing reference strains appropriate for vaccine manufacturing.
- Reviewing antiviral drug and pandemic vaccine supply issues.
- Evaluating and issuing EUAs when appropriate.
- Monitoring vaccine adverse events.
- Monitoring antiviral drug adverse events.
- Maintaining close communication with drug and vaccine manufacturers.
- Evaluating investigational new drug (IND) applications and investigational device exemptions (IDEs) for medical products that diagnose, treat, prevent, or mitigate influenza.
• Evaluating new manufacturing sites and processes for antiviral drugs.
• Making necessary changes in prescribing and patient information, including dosing, target populations, and other directions for use for antiviral drugs and pandemic vaccines based on research and adverse events.
• Evaluating long-term stability of stockpiled antiviral drugs for purposes of shelf-life extension.
• Monitoring to protect against the distribution of counterfeit antiviral drugs and pandemic vaccines.

**HHS Pandemic Influenza Implementation Plan**

The *HHS Pandemic Influenza Implementation Plan* (November 2006) provides a roadmap for the Department’s pandemic preparedness and response. It outlines specific steps to implement the actions identified in the *HHS Pandemic Influenza Plan, Part 1 – Strategic Plan*.  

**F.3 FDA SUPPORTING DOCUMENTATION**

**FDA OPDIV Section to the HHS Pandemic Influenza Operational Plan**

The FDA OPDIV section to the *HHS Pandemic Influenza Operational Plan* (November 21, 2006) includes essential functions that FDA Centers and Offices must continue to perform to ensure COOP in the event of an influenza pandemic.

**Center/Office Emergency Plans and Procedures**

The following plan supports implementation of this *Pandemic Influenza Annex to the FDA EOP*:

• **OC/OO/Office of Financial Operations (OFO)/OFM – Business Continuity Plan (June 2013)**
Appendix A:
Protection of Workers: Guidance on Preparing the Workplace and Workers for an Influenza Pandemic

Introduction
An influenza pandemic occurrence will cause large numbers of employees to be absent from work because of illness, because they may have to care for ill family members, because schools or daycare centers are closed, or because they are afraid to come to work.

It is estimated that during a period of peak influenza pandemic illness, approximately 30 to 40 percent of employees working in a particular office or organization could be absent. An outbreak in a particular geographic area could last from 6 to 8 weeks. There is little way of knowing when an influenza pandemic will occur or how severe it will be in the population.

Planning for an influenza pandemic should assume a worst-case scenario that would be characterized by a high level of illness and absenteeism, social disruption, and economic loss in the public, private, and nonprofit sectors of the economy. The level of risk experienced by employees in the various FDA Centers and Offices will depend on the nature of their occupations and the degree to which they have occupational exposure to people potentially infected with the pandemic influenza virus.

Action Steps
Expanding teleworking by employees and “social distancing” are two actions that can be taken by managers to facilitate continuation of an agency’s essential functions despite high workforce absenteeism. Social distancing refers to employee behavior to reduce the frequency, proximity, and duration of contact between people (both employees and customers) to reduce the chances of spreading influenza from person-to-person.

Spread of Influenza
It is important that employees understand how influenza can be spread among people. It is spread primarily through large droplets (droplet transmission) that directly contact the nose, mouth, or eyes. Droplets are produced when infected people cough, sneeze, or talk. This sends relatively large infectious droplets and very small sprays (aerosols) into the nearby air and into contact with other people. The large droplets can only travel around 6 feet. Consequently, people should limit close contact with others to a distance no closer than 6 feet. A second way that influenza is transmitted is by individuals touching objects that have been contaminated with influenza viruses and then transferring the infected material from their hands to their nose, mouth, or eyes. A third way of transmitting influenza is through the spread of very small infectious particles (aerosols) that travel in the air. Employees need to understand these three principal means of transmission of influenza so that they can take steps, to the maximum extent possible, to protect themselves against infection.

Occupational Safety and Health Administration Guidance
The U.S. Department of Labor (DOL) Occupational Safety and Health Administration (OSHA) has issued a document entitled Guidance on Preparing Workplaces for an Influenza Pandemic (OSHA 3327-02N 2007 and updated version 3227-05R 2009), which provides detailed, but strictly advisory, guidance for employers on preparing their workplaces to minimize the impact of an influenza pandemic on their workers and operations. This guidance will help employers and employees identify the pandemic influenza risk levels in their individual workplace settings. OSHA identifies appropriate control measures for pandemic influenza—these measures include good hygiene, cough etiquette, social distancing, the use of PPE, and staying home from work when ill.
Although the work of the FDA Centers and Offices differs from other USG agencies and among themselves, there are many pandemic preparedness recommendations in the OSHA guidance publication that are applicable to all FDA elements. They include:

- Be aware of and review other USG, State, and local health department pandemic influenza plans; incorporate appropriate actions from these plans into workplace disaster plans.
- Prepare and plan for operations with a reduced workforce.
- Work with your suppliers to ensure you can continue to operate and provide services.
- Develop a sick leave policy that does not penalize sick employees, thereby encouraging employees who have influenza-related symptoms to stay home so they do not infect other employees; recognize that employees with ill family members may need to stay home to care for them.
- Identify possible exposure and health risks to your employees: Are your employees expected to have a lot of contact with the general public?
- Minimize exposure to fellow employees or the public.
- Identify business-essential functions and people required to sustain business-necessary functions and operations; prepare to cross-train or develop ways to function in the absence of these positions. It is recommended that employers train three or more employees to be able to sustain business-necessary functions and operations.
- Plan for downsizing services, but also anticipate any scenario that may require a surge in your services.
- Some employees will have individual risk factors that should be considered by employers as they plan how the organization will respond to a potential pandemic (e.g., immuno-compromised individuals and pregnant women).
- Stockpile items such as soap, tissue, hand sanitizer, cleaning supplies, and recommended PPE. When stockpiling items, be aware of each product’s shelf-life and storage conditions, and incorporate product rotation into your stockpile management program (Guidance on Preparing Workplaces for an Influenza Pandemic, pp.13-15).

OSHA’s Guidance on Preparing Workplaces for Pandemic Influenza notes that the two most effective ways of protecting employees from pandemic influenza depend upon emphasizing proper hygiene (i.e., disinfecting hands and surfaces) and practicing social distancing as previously described. OSHA describes a “hierarchy of controls” as the most effective way of dealing with a workplace hazard. This concept places a high priority on intervention strategies by management and on attempts to remove the hazard from the workplace rather than to rely primarily on the employees to reduce their exposure.

Depending on the nature of the work of each FDA Center and Office, the strategies for work practice and engineering controls, administrative controls, work practices, and PPE will differ in both cost and effectiveness.

**CDC Guidance**

The CDC has issued a report entitled *Community Mitigation Guidelines to Prevent Pandemic Influenza — United States, 2017*, and supporting technical documents which provides planning guidance for State,
local, tribal, and territorial communities. This planning focuses on several measures other than vaccination and drug treatment that might be useful during an influenza pandemic to reduce its harm. The guidelines provide evidence-based recommendations on the use of nonpharmaceutical interventions (NPIs) in mitigating the effects of pandemic influenza. The pandemic mitigation interventions described in the documents include:

**Personal NPI**

- Personal Protective Measures for Everyday Use: Voluntary Home Isolation, Respiratory Etiquette, and Hand Hygiene
- Personal Protective Measures Reserved for Pandemics: Voluntary Home Quarantine and Use of Face Masks in Community Settings

**Community NPI**

- School Closures and Dismissals
- Social Distancing Measures for Schools, Workplaces, and Mass Gatherings

**Environmental NPI**

- Environmental Surface Cleaning Measures or Schools, Workplaces,

**Hierarchy of Controls**

OSHA defines work practice controls as “procedures for safe and proper work that are used to reduce the duration, frequency, or intensity of exposure to a hazard.” Managers and supervisors should solicit suggestions from employees on ways to implement safe work practice controls. Engineering controls are preferred over all others because “they make permanent changes to reduce exposure to hazards and do not rely on employee or customer behavior.” These types of controls involve making permanent changes to the work environment to reduce work-related hazards.

Administrative controls include “controlling employees’ exposure by scheduling their work tasks in ways that minimize their exposure levels.” Examples of administrative controls include discontinuation of non-essential travel to locations with high illness transmission rates, implementing work practices such as email, websites, and teleconferences that minimize face-to-face contact between employees, and adopting flexible work arrangements, such as telecommuting or flexible work hours, to reduce the number of employees who must be at work at one time or in one specific location.

PPE is considered appropriate for employees during certain types of exposures, but is no replacement for other types of prevention interventions, such as engineering controls, cough etiquette, and hand hygiene. Examples of the types of PPE appropriate for a pandemic influenza include: gloves, goggles, face shields, and respiratory protection devices.

Each of the above types of work practices and controls may be appropriate for implementing in FDA Centers and Offices. However, there is no single list of these four prevention strategies that would necessarily be appropriate for implementing in each FDA organization. Rather, it is incumbent upon the director of each Center and Office to engage the employees and supervisors in their unit to examine how the work of their organization can be modified to provide maximum protection for the staff should an influenza pandemic occur.

All actions taken by FDA managers and supervisors must be consistent with and not conflict with other Federal, State, and local health department pandemic influenza plans. As appropriate, portions of these other plans may be incorporated into individual pandemic influenza plans developed for individual FDA Centers and Offices.
HHS, Office of Personal Management (OPM), and U.S. General Services Administration (GSA) have documents and websites that provide policy and guidance applicable for emergency situations, such as an influenza pandemic. Human capital management, leave, temporary closing of work spaces and treatment of absences, and telework are all functions that may need to be implemented during the course of an influenza pandemic that may impact FDA Centers and Offices. As needed, directors and supervisors should consult the following documents/websites:

- OPM and GSA Policy on Telework
  www.telework.gov

- HHS Telecommuting Program Policy
  http://intranet.hhs.gov/ohr/telework/policy.html

- Interim Pre-pandemic Planning Guidance: Community Strategy for Pandemic Influenza Mitigation in the United States – Early, Targeted, Layered Use of Nonpharmaceutical Interventions
  www.flu.gov/planning-preparedness/community/mitigation.html#