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SUMMARY of CHANGE

AR 40-5
Preventive Medicine

This revision--

- Consolidates AR 40-5, AR 40-26, and AR 40-554.
- Adds Responsibilities for commanders (chaps 1, 9, and 12) and preventive medicine personnel (chaps 1, 2, 4, 5, 6, and).
- Adds information on the Preventive Dentistry Program, community health nursing activities, disease and climatic injury prevention and control, medical examinations, spirometry surveillance, community and family health, nonionizing radiation registry, sanitation, and field preventive medicine (chaps 2, 3, 4, 5, 6, 9, 12, and 14).
- Deletes the appendix on ice manufacture sanitation in AR 40-5.
- Rescinds RCS MED-292 (DA Form 3898-R (Report of Tuberculosis Detection and Control)).
- Adds DD Form 2493-1 (Asbestos Exposure, Part I--Initial Medical Questionnaire).
- Adds DD Form 2493-2 (Asbestos Exposure, Part II-Periodic Medical Questionnaire).
- Adds DA Form 3897-R (Tuberculosis Registry).
- Adds DA Form 5931 (Occupational Health Patient Form).
- Adds DA Form 5932 (USAREUR Occupational Health Form).
- Adds DA Form 5933 (Occupational Health Patient Form-Supplemental).
- Adds DA Form 5934 (Korea Occupational Health Encounter Form).
- Adds DA Poster 40-5 (Lyme Disease Warning).
By Order of the Secretary of the Army:

CARL E. VUONO
General, United States Army
Chief of Staff

Official:

MILTON H. HAMILTON
Administrative Assistant to the Secretary of the Army

History. UPDATE printing of November 1990 published a revision of this publication. This publication has been reorganized to make it compatible with the Army electronic publishing database. No content has been changed.

Summary. This regulation is a consolidation of several regulations that cover the Army's preventive medicine program. It establishes practical measures for the preservation and promotion of health and the prevention of disease and injury. This regulation implements Executive Order 12196 and DOD Instructions 6050.5, 6055.1, 6055.5, and 6055.12.

Applicability. This regulation applies to facilities controlled by the Army and to all elements of the Army. This includes military personnel on active duty; U.S. Army Reserve or Army National Guard personnel on active duty or in drill status; U.S. Military Academy cadets; U.S. Army Reserve Officer Training Corps cadets, when engaged in directed training activities; foreign national military personnel assigned to Army components; and civilian personnel and nonappropriated fund employees who are employed by the Army on a worldwide basis.

Army management control process. This regulation is subject to the requirements of AR 11–2. This regulation contains internal control provisions but does not contain checklists for conducting internal control reviews. These checklists are contained in DA Circular 11–88–7.

Supplementation. Supplementation of this regulation by the principal HQDA officials and major Army commands listed below is permitted. Supplementation is prohibited by all other elements without prior approval of HQDA (SGPS–PSP), 5109 Leesburg Pike, Falls Church, VA 22041–3258. If supplements are issued, one copy of each will be furnished to HQDA (SGPS–PSP), 5109 Leesburg Pike, Falls Church, VA 22041–3258.

Interim changes. Interim changes to this regulation are not official unless they are authenticated by the Administrative Assistant to the Secretary of the Army. Users will destroy interim changes on their expiration dates unless sooner superseded or rescinded.

Suggested Improvements. The proponent agency of this regulation is the Office of The Surgeon General. Users are invited to send comments and suggested improvements on DA Form 2028 (Recommended Changes to Publications and Blank Forms) directly to HQDA (SGPS–PSP), 5109 Leesburg Pike, Falls Church, VA 22041–3258.

Distribution. Distribution of this publication is made in accordance with the requirements on DA Form 12–09–E, block number 2058, intended for command level C for Active Army, Army National Guard, and U.S. Army Reserve (applicable to all Army elements); and command level A for Active Army and Army National Guard and D for U.S. Army Reserve (applicable to medical activities only).
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### Glossary

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RESERVED
Chapter 1

Introduction

1–1. Purpose

This regulation—

a. Explains the Army Preventive Medicine Program.

b. Prescribes a comprehensive disease prevention and environmental enhancement plan of action for the U.S. Army at fixed installations and in support of field forces.

c. Establishes military occupational and environmental health standards.

d. Defines the activities within the Preventive Medicine Program-functional areas.

e. Provides a basic guide for commanders, the installation medical authorities(IMAs), and other interested persons and agencies.

f. Contains policy, guidelines, and procedures.

g. Provides organizational structure guidance.

h. Describes the functions and responsibilities of preventive medicine (PVNTMED) services at the U.S. Army medical department activity(MEDDAC) and U.S. Army medical center (MEDCEN) level.

i. Identifies Department of the Army (DA) occupational safety and health (OSH) standards.

1–2. References

Required and related publications and prescribed and referenced forms are listed in appendix A.

1–3. Explanation of abbreviations and terms

Abbreviations and special terms used in this regulation are explained in the glossary.

1–4. Responsibilities

a. The Assistant Secretary of the Army for Installations and Logistics, in addition to the responsibilities cited in AR 385–10, will—

   (1) Provide executive leadership at the Army Secretariat level to ensure timely compliance with environmental, safety, and occupational health (OH) requirements.

   (2) Establish goals and policies and monitor programs for environmental, safety, and OH.

b. The Surgeon General (TSG) is responsible for the overall development and oversight of DA policies and programs for the Army-wide Preventive Medicine Program, which includes—

   (1) Disease and climatic injury control.

   (2) OH.

   (3) Community and family health.

   (4) Health information and education.

   (5) Nutrition.

   (6) Health hazard assessment (HHA).

   (7) Medical safety.

   (8) Radiation protection.

   (9) Pest and disease vector control.

   (10) Environmental quality.

   (11) Sanitation.

   (12) Environmental laboratory services.

   (13) Design review.

   (14) Field PVNTMED.

   (15) Toxicology.

c. The director, Army Safety, Office of the Army Safety Program, will carry out the responsibilities defined in AR 385–10.

d. The Chief, Preventive and Military Medicine Consultants Division, Office of the Surgeon General (OTSG) will—

   (1) Formulate policies, standards, regulations, and directives to protect and promote health, improve effectiveness, and enhance the environment of Army personnel.

   (2) Exercise staff supervision, program management (including Army Regulation (AR) propensy), and provide consultative services on the Army-wide Preventive Medicine Program described in b above.

   (3) Monitor and act as point of contact (POC) on health and welfare aspects of environmental quality.

   (4) Advise and assist the Army staff in development of DA plans, policies, and regulations on health conservation and control of environmental quality.

   (5) Provide international and interservice representation and liaison with professional organizations, Department of Defense (DOD), and other Federal agencies to exchange data on disease control, health maintenance, and environmental medicine.

   (6) Determine appropriate preventive measures, pharmaceuticals, and biologics for disease control and initiate requests for supply actions to ensure availability.

   (7) Coordinate with the DA Safety Office for compliance with Occupational Safety and Health Act health standards.

   (8) Provide administrative support and staff supervision to the Armed Forces Epidemiological Board and the Armed Forces Pest Management Board(AFPMB).

   (9) Evaluate and approve requests for epidemiology consultant(EPICON) assistance in the study of disease outbreaks.

   (10) Be the OTSG reviewing authority for all environmental documents submitted by DA activities.

   (11) Provide Preventive Medicine Program direction through the U.S. Army Health Services Command (HSC) to U.S. Army Environmental Hygiene Agency (USAEHA) and to U.S. Army Aeromedical Center(DA missions), through U.S. Army Japan to U.S. Army Pacific, Environmental Health Engineering Agency (USAPACEHEA), and through 7th Medical Command for 10th Medical Laboratory.

   (12) Coordinate the mission services of USAEHA with appropriate elements of the DA staff and outside continental United States (OCONUS)medical support organizations (see para 1–8c).

   (13) Coordinate directly for USAEHA services provided in support of the DA-level PVNTMED mission.

   (14) Provide professional advice concerning materiel and facilities requirements.

   (15) Conduct HHA of medical and nonmedical material.

e. All major Army command (MACOM) commanders will establish a formal procedure to respond to the USAEHA, 10th Medical Laboratory, and USAPACEHEA report recommendations involving regulatory compliance. Further, the commanders will monitor compliance, and this procedure must provide for—

   (1) Tracking the corrective actions involving regulatory compliance and target dates for completing planned action.

   (2) Issuing copies of the installation’s responses and planned corrective actions to the report originator (USAEHA, 10th Medical Laboratory, or USAPACEHEA) for review and comment.

   (3) Reporting the status of uncorrected problems identified in USAEHA, 10th Medical Laboratory, or USAPACEHEA reports in annual environmental and OH management reports as prescribed by Headquarters, Department of the Army (HQDA).

   f. The commanding general, HSC will—

   (1) Provide health care services and resources for the Army within the continental United States (CONUS), Alaska, Panama, Puerto Rico, Hawaii, Johnston Island, Guam, and the trust territories of the Pacific.

   (2) Plan, program, and budget resources for the USAEHA.

   (3) Provide command guidance on the priorities, services, and direction of USAEHA.

   g. Commanders at all levels will promote general health and safety and ensure occupational and environmental health within their commands. Commanders will—

   (1) Support the Preventive Medicine Program.

   (2) Provide adequate resources to implement the program.

   (3) Take appropriate actions, based on recommendations of the IMAs, to protect all personnel under their jurisdiction from disease and injury.

   (4) If DA Poster 40–5 (Lyme Disease Warning) is used, follow guidance in paragraph 10–18c.

   h. Commanders of dental activities (DENTACs) will—

   (1) Implement and monitor the Army Preventive Dentistry Program per AR 40–35.
(2) Forward a copy of the preventive dentistry report to the appropriate MACOM surgeon.

i. The IMAs are responsible to commanders for the following:
(1) Establishing and operating an effective Preventive Medicine Program. The program will be supported by adequate—
(a) Personnel,
(b) Funding,
(c) Office and laboratory space,
(d) Equipment and supplies,
(e) Transportation and communications.
(2) Recommending solutions for all PVNTMED problems.
(3) Providing PVNTMED guidance based on the functional areas described in this regulation.

j. Heads of installation civilian personnel offices will take the following actions to assist medical personnel with the medical evaluation:
(1) Identify employees expected to be absent from work for 2 weeks or more.
(2) Provide Army medical personnel with Department of Labor (DOL) Forms CA–16 (Authorization for Examination and/or Treatment) and CA–17 (Duty Status Report) (or equivalent medical documentation) for completion by the treating physician for those employees identified.
(3) Make arrangements with employees for examinations whenever necessary.

k. The commander, USAEHA will—
(1) Provide worldwide support of PVNTMED programs for the Army through consultations, supportive services, investigations, and training in the areas of environmental quality, occupational and environmental health, toxicology, disease prevention, surveillance and control, radiation and environmental sciences, pest management, and laboratory services.
(2) Evaluate the responses to recommendations and resolve situations with MACOMs where responses to recommended corrective actions to USAEHA reports are considered unsatisfactory. When the MACOM and USAEHA cannot agree on proposed corrective actions, the matter with all associated correspondence will be referred to USAEHA through HSC, or the appropriate OCONUS medical support organization, to HQDA (SGPS–PSP), 5109 Leesburg Pike, Falls Church, VA 22041–3258. OTSG will coordinate with appropriate Army staff and provide a resolution to the MACOM.
(3) Review proposed environmental, safety, and health standards or standards criteria documents published for comment by regulatory agencies and consensus standard organizations. The commander also will provide written technical comment regarding content, feasibility of implementation, and applicability to Army operations. In addition, the technical input of the U.S. Army Safety Center and other DA organizations will be solicited as necessary to facilitate such review.
(4) Conduct the Army Preventive Medicine Residency Training Program in occupational medicine to meet accreditation requirements of the American Council on Graduate Medical Education and approval requirements of the American Osteopathic Association.

I. Commanders, 10th Medical Laboratory and USAPACHEA will—
(1) Provide theaterwide support of PVNTMED programs for the Army through consultations, supportive services, investigations, and training in the areas of environmental quality, occupational and environmental quality, occupational and environmental health, toxicology, disease prevention, surveillance and control, radiation and environmental sciences, pest management, and laboratory services, as staffing permits.
(2) Evaluate the responses to recommendations and resolve situations with MACOMs where responses to recommended correction actions to 10th Medical Laboratory or USAPACHEA reports are considered unsatisfactory. When the MACOM and 10th Medical Laboratory or USAPACHEA cannot agree on proposed corrective actions, the matter with all associated correspondence will be forwarded by 10th Medical Laboratory or USAPACHEA to HQDA (SGPS–PSP), 5109 Leesburg Pike, Falls Church, VA 22041–3258.

OTSG will coordinate with appropriate Army Staff and provide a resolution to the MACOM.

m. Managers and supervisors are responsible for—
(1) Keeping informed of OH hazards and requirements in activities under their control.
(2) Training employees in appropriate health and safety practices.
(3) Enforcing the use of protective clothing and equipment.
(4) Providing the civilian personnel office with health and safety information necessary for effective job classification and placement actions.

1–5. Program concept
DA policy is to conserve the fighting strength by controlling preventable disease and injury through command-oriented occupational, environmental, and personal protection programs. The individual’s role in maintaining his or her own health and fitness will be emphasized.

1–6. Liaison
a. Liaison will be established and maintained at all organizational levels with medical departments of other military services, and appropriate representatives of Federal, State, and local health and environmental protection authorities (AR 200–1).

b. Participation on Armed Forces disciplinary control boards and liaison with representatives of civil agencies concerned with health and welfare are prescribed in AR 190–24/MCO 1620.2/BUPERINST 1620.4/AFR 125–11/COMDDINST 1620.1.

1–7. Recordkeeping
AR 25–400–2 establishes the Modern Army Recordkeeping System (MARKS). This system reorganized the files listed in The Army Functional File System by identifying each file by the number of the directive prescribing that those records be created, maintained, and used. Therefore, records required by this regulation should be filed under the file number 40–5. Refer to AR 25–400–2, appendixes B, C, or D for further guidance.

1–6. Technical assistance
a. Commanders and IMAs at all levels may request technical assistance in matters pertaining to the Preventive Medicine Program through command channels.

b. CONUS requests should be addressed through the MACOM command channels of the activity requesting services to the Commander, USAEHA, Aberdeen Proving Ground, MD 21010–5422, with a copy furnished to Commander, HSC, ATTN: HSCL–P, Fort Sam Houston, TX 78234–6000.

c. OCONUS requests from the—
(1) U.S. Army, Europe (USAREUR) and Seventh Army areas of responsibility will be forwarded to Commander, 7th Medical Command, ATTN: AEMCL–PM, APO New York 09102.

(2) Pacific geographic areas of responsibility will be forwarded to Commander, USAPACHEA—Sagami, APO San Francisco 96343.

(3) U.S. Army South areas of responsibility will be forwarded to Commander, U.S. Army South, ATTN: SOMD, APO Miami 34004.

(4) If 7th Medical Command, USAPACHEA, or U.S. Army South cannot provide the requested services, the requests will be forwarded to HQDA(SGPS–PSP), 5109 Leesburg Pike, Falls Church, VA 22041–3258.

Chapter 2
Army Preventive Medicine Program

Section I
Background

2–1. General
The Preventive Medicine Program is a comprehensive program, ranging from simple field sanitation procedures to extensive and
complicated monitoring techniques necessary to protect the health and environment of Army personnel. The program is designed to promote and maintain the fighting force at maximum effective strength and to maintain the physical well-being of all personnel for whom the Army is responsible.

2–2. Functional areas
The PVNTMED functional areas are as follows:

a. Disease and climatic injury prevention and control. This includes communicable disease control, chronic disease prevention, hospital infection control, nutrition, and prevention of injury related to heat, cold, altitude, and other environmental elements. (See chap 4.)

b. OH. This includes activities designed to focus on the person and his or her occupational environment with the goal of providing a safe and healthful workplace. These activities encompass (refer to chap 5)—

(1) Assessment of OH hazards associated with materiel, raw materials, by-products, processes, and practices inherent to the work environment.

(2) Establishment of criteria for the evaluation and control of occupational exposures.

(3) Determination of requirements for and of provisions of medical surveillance and worker health education.

(4) Application of epidemiological principles in evaluating the work environment.

(5) Recognition, evaluation, and prescription of methods to control environmental factors that may adversely affect employee health or well-being.

c. Community and family health. This includes those activities that promote family health and the health of service members within the military community. Also included are those services and activities that establish an interface between the medical treatment facility (MTF) and the community, as well as health programs designed to reach specific individuals or groups at the community level. (See chap 6.)

d. Health information and education. This includes health information programs for the general community and health education programs for the individual soldier. The orientation will be toward behavioral modification to improve health and limit disability by encouraging individual responsibility. (See chap 7.)

e. Nutrition. This includes the development of nutrition standards and policy regarding the soldier’s diet both in garrison and in field settings (combat rations). Standards are published in AR 40–25/NAVMEDCOMINST 10110.1/AFR 100–95 and periodically revised as required.

f. HHA. This includes activities to support the Manpower Personnel Integration Program by identifying potential health hazards associated with the life-cycle management of weapons, equipment, training devices, and materiel systems and by recommending appropriate efforts to either eliminate or control such hazards. See AR 40–10 for the following:

(1) Objectives and policies of the Army’s Health Hazard Assessment Program in support of the Army materiel acquisition decision process (MADP).

(2) Specific responsibilities of the Army staff, MACOMs, testers and evaluators, and developers for HHAs.

(3) Specific procedures, including the coordination of HHA with system safety and human factors engineering portions of the MADP.

(4) Procedures to identify and eliminate or control health hazards associated with MADP, including preparation of the Health Hazard Assessment Report (Requirement Control Symbol (RCS) MED-388).

g. Medical safety. This includes the Army Medical Department (AMEDD) unit safety program, hospital safety, and medical safety systems for both garrison and field operations. (See chap 8.)

h. Radiation protection. This encompasses both ionizing and non-ionizing radiation, to include licensing and authorizations, personal protective measures, radiation detection and measuring equipment, control of radiation sources, radiation shielding, and operational surveillance. (See chap 9.)

i. Pest and disease vector prevention and control. This includes prevention and control of disease vectors and animal reservoirs, integrated pest management (IPM) operations and research, disease vector surveillance, pest quarantine, and retrograde cargo inspection and treatment. A close working relationship with the AFPMB is maintained to ensure DA input into DOD pest management programs. (See chap 10.)

j. Environmental quality. This includes all AMEDD subprograms in support of the Army environmental program that has as its purpose the protection and preservation of environmental quality related to the health and welfare of DA personnel. (See chap 11.)

k. Sanitation. This includes subprograms to improve environmental conditions for the maintenance of health. (See chap 12.)

l. Environmental laboratory services. This includes laboratory services required to support all PVNTMED programs that are provided by the installation, USAEHA, and other regional laboratories. The level of support required is determined by regulatory agency guidance and DA directives. (See chap 13.)

m. Design review. This includes review of the health aspects of drawings, plans, and related technical documents for projects such as food service, troop housing, MTFs, and many industrial manufacturing and maintenance facilities. PVNTMED personnel participate in this program at the installation, regional, MACOM, and DA level.

n. Field PVNTMED. This includes training requirements and operational responsibilities for units deployable to the field. (See chap 14.)

o. Toxicology. This includes support to the Preventive Medicine Program by means of—

(1) Toxicological assessments and laboratory evaluations of potentially hazardous materials.

(2) Toxicity clearances and health risk assessments, as appropriate. USAEHA may be required to conduct animal testing, toxicity studies, and literature reviews to support this program.

Section II
PVNTMED Levels of Support and Special Resources

2–3. General
The Preventive Medicine Program is organized and staffed on a decentralized basis. PVNTMED services are provided on the following three levels:

a. First level. Local PVNTMED services are provided on the basis of tables of organization and equipment (TOE) and tables of distribution and allowances (TDA). It also includes services provided by PVNTMED TOE teams and MEDDACs when assigned in a direct support role.

b. Second level. PVNTMED officers assigned to CONUS MEDCENs and the 7th Medical Command (USAREUR) and the 7th Medical Command (USAREUR) and the 7th Medical Command (USAREUR) Regional Laboratory have as additional responsibilities for units deployable to the field.

(1) Infectious diseases.

(2) Occupational diseases (in collaboration with USAEHA).

(3) Chronic diseases and nonbattle injuries.

(4) Public health aspects of disaster relief operations.

(5) Design of medical studies.

(6) Design of medical studies.
2–5. Intercommand relationships

Relationships regarding PVNTMED services crossing MACOM lines will be addressed in a memorandum of understanding.

Section III

PVNTMED Personnel

2–6. General

Maximum use will be made of the professional capabilities of PVNTMED personnel in direct support of the Preventive Medicine Program. Additional duties will not be allowed to interfere with the performance of professional duties outlined in AR 611–101.

2–7. Activities of the Chief, PVNTMED service

a. The chief will establish and direct the Preventive Medicine Program for the supported health service area as described in this regulation.

b. The chief or his or her designee should—

(1) Serve as consultant and provide liaison to the installation commander, his or her staff, and tenant activities in PVNTMED.

(2) Establish and maintain liaison with appropriate Federal, State, and local health authorities.

c. The chief may be the AMEDD representative on installation boards, councils, and committees.

d. When a residency-trained PVNTMED officer or occupational medicine officer (area of concentration (AOC) 60C or 60D) is assigned, he or she will direct the program. Otherwise, the chief will be an AMEDD officer, usually the senior officer, assigned to the PVNTMED service.

Chapter 3

Reports

Section I

Special Telegraphic Reports (RCS MED–16)

3–1. General

Submission of the RCS MED–16 requires liaison among PVNTMED personnel, the patient administration division of the MTF, and the medical staff (AR 40–400, chap 6).

3–2. Reporting guidance

Reporting requirements for the special telegraphic reports of selected conditions, reportable outbreaks, and reportable deaths are outlined in AR 40–400.

Section II

Command Health Reports (RCS MED–3)

3–3. General

This section establishes procedures for the periodic reporting of all matters pertaining to the health of the command. Command health reports (CHRs) are forwarded through command channels and are designed to—

a. Inform commanders of health conditions within their commands and recommend measures for improvement.

b. Provide commanders an opportunity to record actions taken for improvement and to inform higher headquarters of support required to implement recommendations.

c. Provide information on unsolved problems, new developments, and other matters relating to command health effectiveness.

d. Provide data for the periodic assessment of Preventive Medicine Program effectiveness.

e. Serve as feeder reports for preparation of a consolidated CHR.

3–4. Preparing agencies

a. The CHR for fixed installations will be prepared and signed by the director, health services (DHS) and addressed to the installation commander.

b. The CHR for battalion-sized or larger units will be prepared and signed by the unit surgeon and addressed to the unit commander.

c. The CHR for units or installations without unit surgeons or DHSs will be included in the CHR of the unit surgeon or DHS rendering primary medical services. The CHR will be addressed to the commander of the supported installation.

d. For MACOMs and Army components of unified commands as listed in AR 10–5, chapter 3, a consolidated CHR will be prepared.

e. Installations or activities such as separate recruiting offices, military entrance processing stations, Reserve Officers’ Training Corps units, and other off-post activities located at civilian facilities not under control of Government-owned, contractor-operated installations are exempt from CHR preparation requirements of this regulation.

3–5. Frequency

a. Installation and unit CHRs will be prepared the last day of each calendar month.

b. Consolidated CHRs for MACOM and Army components of unified commands will be prepared the last day of each calendar quarter.

3–6. Due dates

a. Commanders of MACOMs will establish due dates for feeder reports.

b. Consolidated reports will be dispatched quarterly by the MACOM to HQDA (SGPS–PSP), 5109 Leesburg Pike, Falls Church, VA 22041–3258, with a copy furnished to Commander, HSC, ATTN: HSCL–P, Fort Sam Houston, TX 78234–6000, to be received not later than 30 working days following the end of the report period. Copies of OCONUS consolidated reports will not be furnished to the HSC, with the exception of those relating to the U.S. Army South.

3–7. Command routing

a. Commanders will endorse the CHRs, as originally submitted by the IMA through command channels, and will note approval or disapproval and actions taken to improve conditions and correct reported deficiencies.

b. Commanders of major Army field commands and Army components of unified commands will establish internal routing and consolidation procedures for CHRs within their command. The reports will be screened and the data used as a basis for comments to be submitted to the next higher command.

c. An information copy of the CHR prepared by HSC MTFs and activities will be forwarded to the Commander, HSC, ATTN: HSCL–P, Fort Sam Houston, TX 78234–6000.

3–8. Preparation instructions

The CHR will be prepared by the unit surgeon and/or DHS and will include current information on PVNTMED matters. Outstanding accomplishments, new developments, and trends will be recorded as well as unsatisfactory conditions and recommended corrective measures. As a rule, unsatisfactory conditions that are identified and are readily correctable locally need not be reported. Unsatisfactory health conditions that persist and cannot be corrected locally because of the lack of resources must be reported to higher headquarters. If unsatisfactory health conditions are reported, suitable
corrective measures will be recommended at the conclusion of the CHR. The CHR is not intended to convey routine or repetitious information regarding satisfactory conditions. If no comment is required under a particular heading, the heading will be omitted; a negative comment is not required. To provide uniformity, the following headings and paragraph designations will be used (when applicable):

a. Health of the command.
   (1) Personal hygiene.
      (a) Sanitary discipline.
      (b) Condition and adequacy of personal protective equipment (PPE) and clothing.
   (2) Status of training of unit field sanitation teams (para 14–3b).
   (3) Disease.
      (a) If there has been an increased incidence of any disease of military importance among military or civilian personnel, the situation will be reported from an epidemiological viewpoint (such as diagnosis (presumptive or confirmed), scope, population distribution, immunization status (when applicable), control measures, and course of the outbreak).
      (b) Note the occurrence of any unusually severe reactions to immunizations.
      (c) Nosocomial infection rate will be reported as specified in paragraph 4–11 (by those hospitals conducting total surveillance).
   (4) Injuries.
      (a) Heat and cold injury. Discuss cases from an epidemiological viewpoint, noting trends, host and environmental factors, and status of training and education.
      (b) Other injuries. Discuss injuries that are significant or unusual, such as trainees’ stress fractures and soldiers’ eye injuries.

b. Occupational health.
   (1) Status of completion or revision of the Health Hazard Information Module (HHIM), which is part of the Occupational Health Management Information System (OHMIS).
   (2) Design review activities.
   (3) Significant trends in incidence of occupational illness or injury.
   (4) Status of Occupational Safety and Health Administration (OSHA) abatement plans of occupational hazards or any other significant problems.

c. Environmental sanitation.
   (1) Water supply, including sources and method of treatment, fluoridation, adequacy of quantity and quality, and changes or additions to the water supply system. Variations from standards will be reported under AR 420–46, paragraph 5d.
   (2) Liquid waste disposal, including—
      (a) Type of system and method of treatment.
      (b) Size and type of receiving streams.
      (c) Possible health hazards and public nuisances.
      (d) Changes or additions to the system.
   (3) Industrial-type waste disposal originating on military installations and incident to military operations (including source, amount, type, and method of treatment or disposal).
   (4) Solid waste management, including—
      (a) Method and place of disposal.
      (b) Truck washing facilities.
      (c) Interim storage facilities.
      (d) The use of volume reduction equipment.
      (e) Specialized collection vehicles.
   (5) Housing and other buildings, including—
      (a) Type and condition.
      (b) Floor space available per person.
      (c) Ventilation.
      (d) Heating.
      (e) Lighting.
      (f) General cleanliness.
      (g) Adequacy of latrine and laundry facilities, as applicable.
      (h) Food service facilities, including—
         (a) Type, condition, and adequacy of food service buildings.
         (b) Equipment (to include adequacy of refrigeration, dishwashing facilities, latrine facilities, insect and rodent control, and waste disposal).
   (7) Specific sanitation problems related to—
      (a) The operation of recreational facilities.
      (b) Child development centers.
      (c) Barber and beauty shops.
      (d) Ice plants.
      (e) Mobile home parks.
      (f) Nuisances adjacent to the installation.
   (8) Sanitation problems unique to health care facilities, including—
      (a) Collection and disposal of infectious waste.
      (b) Housekeeping practices.
      (c) Use of disinfectants and sterilants.
      (d) Ventilation deficiencies.
      (e) Significant pest infestations (particularly in patient-sensitive areas) and efforts to control such infestations.
   (9) Swimming pools and bathing beaches (fresh or salt water), including—
      (a) Types and construction of pools.
      (b) Sanitary and bacteriological quality of water.
      (c) Adequacy of safety features.
      (d) Bathhouse and shower construction.
      (e) Furnishing and operation.
   d. Environmental enhancement.
      (1) Visits and inspections by Federal, State, or local environmental pollution control agencies with recommendations and corrective action.
      (2) Complaints received alleging pollution from sources under Army jurisdiction, and corrective action.
   e. Pest management and pesticide monitoring. Concerns include—
      (1) Significant vector-borne disease occurrences.
      (2) Effectiveness of pest surveillance and control programs to include availability and problems with pesticides or pesticide dispersal equipment.
      (3) Environmental incidents resulting from the use, storage, or disposal of pesticides.
      (4) Confirmed or suspected health-related problems associated with occupational or other exposure to pesticides.
      (1) Significant observations on troop nutritional status and adequacy of the diet.
      (2) Special actions taken concerning nutritional surveys and their findings, changes in ration components with reasons, and observations regarding the Army weight control program.
   g. Community health nursing. The community health nurse (CHN) will report pertinent information concerning specific programs and additions and deletions. Problem areas, including proposed solutions and conditions that adversely affect the health of the active duty military and their family members, and retirees and their family members, who will be reported.
   h. Liaison activities. Indicate liaison activities with other military and civilian health authorities.
      i. PVNTMED staffing problems. Discuss all persistent staffing problems that interfere with attainment of PVNTMED goals.
      j. New and improved PVNTMED measures. Reporting will be limited to measures not reported elsewhere and include a brief description of measures, use, and effectiveness.
   k. Veterinary data.
      (1) Information concerning control of animal diseases communicable to man will consider—
         (a) The origin and progress of a disease.
         (b) Scope and results of any tests.
         (c) Methods of treatment.
         (d) The nature and efficiency of quarantine (if applicable).
         (e) The number of cases and disposition.
      (2) The report will also include information on unusual conditions affecting food sources, storage and handling facilities, and food items that may endanger the health of the command.
   l. Preventive Psychiatry Program. Include the adequacy of the
Preventive Psychiatry Program to maintain and improve command psychological effectiveness.

m. Other. Include subjects not covered under other headings.

n. Recommendations. Recommendations will be made for the correction of deficiencies noted and for improvement of existing or the adoption of new preventive and sanitary measures.

a. Supporting material. Supporting material required to clarify the narrative report will be appropriately identified, cross-referenced, and included as an annex to the report.

3–9. Special command health notification
Notification concerning special or potentially serious health problems will be transmitted telephonically, followed within 72 hours by a written report, by the IMA to the unit and installation commanders. The purpose is to inform the commander concerning serious sanitary deficiencies, environmental or occupational hazards, potential epidemic conditions, or other serious situations that may affect the health of the command. The report will include recommendations and action taken. The installation commander will forward information copies through command channels to HQDA (SGPS–SPS), 5109 Leesburg Pike, Falls Church, VA 22041–3258, and Commander, HSC, ATTN: HSCL–P, Fort Sam Houston, TX 78234–6000.

Section III
DA Form 3076 (Army Occupational Health Report) (RCS MED–20)

3–10. General
This report provides essential information for TSG to discharge staff responsibilities for planning, directing, and supervising health services in the DA Occupational Health Program as required by AR 10–5.

3–11. Preparing agencies
All Army MTFs responsible for providing OH services for civilian or active duty military personnel as authorized in chapter 5 will prepare this report on a biannual basis.

3–12. Preparation instructions
Instructions for preparation of this report and DA Form 3075 (Occupational Health Daily Log) are as follows:

a. Purpose.

(1) DA Form 3076 provides specific data for use by installation and MEDDAC personnel, and higher headquarters for definitive analysis and review of the Occupational Health Program. This tool should aid in evaluating incidence and trends of occupational illnesses and injuries, the extent and effectiveness of other health maintenance activities, and staffing and related needs and resources.

(2) A local system using the DA Form 3075 report and other appropriate reports will be used by the Occupational Health Program coordinator in collaboration with the patient administration officer and chief, department of primary care and community medicine (DPCCM), as applicable, to record required data on a daily and monthly basis. This should include follow-up and/or supportive information (such as, explanatory descriptions of occupational illnesses, medical surveillance evidence of overexposure to hazards, lost-time eye injuries, and new cases of noise-induced hearing loss). Planning for special one-time activities, such as a disease screening program, should include a system to record all required reporting data.

(3) Occupational injury and illness data will be coordinated with the safety officer to assure complete and accurate recording by both the safety office and the MTF.

(4) Special telegraphic reports of occupational illness will be submitted per AR 40–400.

b. Preparing agencies.

(1) All Army MTFs responsible for providing OH services for civilians or active duty military personnel as authorized in chapter 5 will maintain a DA Form 3075 and will prepare a biannual DA Form 3076. This does not include Army MTFs located at and providing OH services primarily for non-DA(such as the Defense Logistics Agency (DLA)) installations.

(2) The DA Form 3076 will be completed by the medical officer or physician in charge of the Occupational Health Program. If there is no full-time medical officer or civilian physician, the person most knowledgeable about the program (normally the senior OH nurse) will complete the report. All data will be based on DA Form 3075 and other records normally kept by the MTFs providing OH services or by other installation activities, such as the civilian or military personnel office, safety office, audiology clinic, emergency clinic, etc.

c. Consolidated reports.

(1) Where there is more than one MTF at an installation providing OH services for civilian employees or active duty military personnel, each will prepare a separate DA Form 3076. (Examples include troop medical clinics, OH clinics, and outpatient clinics.) A consolidated report for the installation will be submitted with a copy of the DA Form 3076 report for each MTF.

(2) The person responsible for preparing the consolidated report will develop a procedure to assure all required data are collected and reported by all MTF activities providing OH services. This will include—

(a) Instructing all supporting activities regarding reporting requirements and assisting them as necessary.

(b) Providing the supporting activities with copies of DA Forms 3075 and 3076 and other forms, when pertinent, for data collection.

(c) Coordinating with the civilian and military personnel offices, safety officer, radiation protection officer (RPO), audiolist, etc., to obtain essential data.

(3) Except where pertinent or feasible, the MTF feeder reports to the consolidated report are not required. For example, staffing for the troop medical clinic, emergency room, outpatient clinic, etc., will not ordinarily be counted (1) above).

(4) The chief, PVNTMED will review the completed DA Form 3076 with the PVNTMED and OH staff and, where pertinent, with department or clinic staff to assure completeness and accuracy of the report and to determine needs for changes or revisions in the OH services and/or procedures.

d. Reporting period and routing. The DA Form 3076 will be prepared semiannually and dispatched no later than 17 working days following the last day of June and December.

(1) CONUS. MTFs will forward the DA Form 3076 as follows:

(a) The original and one copy through the appropriate command, MEDCEN or MEDDAC to the Commander. USAEHA, ATTN: HSHBOM, Aberdeen Proving Ground, MD 210105422.

(b) One copy to HQDA (SGPS–SPS), 5109 Leesburg Pike, Falls Church, VA 220413258.

(c) One copy to the Commander, HSC, ATTN: HSCLP, Fort Sam Houston, TX 782346000.

(d) One copy through command channels to the surgeon of the appropriate MACOM. (The MACOM surgeon in turn will review the DA Form 3076 and forward specific recommendations or comments to HQDA (SGPS–SPS), 5109 Leesburg Pike, Falls Church, VA 220413258 with one copy to Commander, USAEHA, ATTN: HSHBOM, Aberdeen Proving Ground, MD 210105422.)

(2) OCONUS. MTFs will submit the DA Form 3076 through command channels to HQDA (SGPS–SPS), 5109 Leesburg Pike, Falls Church, VA 220413258, with one copy to the Commander, USAEHA, ATTN: HSHBOM, Aberdeen Proving Ground, MD 210105422.

e. Preparation instructions. Data line items on DA Forms 3075 and 3076 correspond, and the instructions for the two forms are the same.

(1) Support data. Items 1 through 12, DA Form 3076, contain data reported for eligible served and assigned staff and will be based on status as of the last day of the report period. The unnumbered items on DA Form 3075 are self-explanatory.

(2) Injury and illness. Items 13 through 26 are to be used for recording occupational fatalities, injuries, and illnesses. Nonoccupational illness and injury visits are to be recorded in item 27.
(3) **Elective health programs.** Items 28 through 35 include services provided that are not job-related such as light duty workers receiving audiograms and/or vision screening who are not exposed to noise and/or eye hazards at work.

(4) **OH services.** Items 36 through 56 on DA Form 3076 and items 37 through 41, 43, 45 through 48, 50 through 52, and 54 through 56 on DA Form 3075 contain data about job-related health services.

(5) **Remarks.** Attach additional sheets as necessary to provide required data noted above. Also, include pertinent information regarding staffing, staff training, specific OH problems or accomplishments, epidemiological data, and so forth.

Section IV
DA Form 3761(Army Health Nursing Activities)(RCS MED-371)

3–13. General
DA Form 3761 and accompanying narrative provides essential information regarding the Community Health Nursing Program. DA Form 3761 is used to plan and evaluate the overall nursing program and specific nursing activities within the program. It is also used as a feeder report for other MEDDAC and MEDCEN required reports.

3–14. Preparing agencies
Each community health nursing section of a PVNTMED service will prepare the DA Form 3761 on a monthly basis.

3–15. Preparation instructions
Instructions for preparing DA Form 3761 are found in AR 40-407, paragraph 7-5.

Chapter 4
Disease and Climatic Injury Prevention and Control

Section I
Disease Prevention and Control

4–1. General
a. Epidemic potentials include those diseases and injuries that can seriously compromise the ability of a military unit to carry out its mission. Preventive measures are essential. Exercise of command authority based on sound medical recommendations, troop discipline, and provision of PVNTMED services in both garrison and field settings is critical.

b. Disease conditions of greatest epidemic potential include, but are not limited to: Acute respiratory diseases, diarrheal diseases, hepatitis, and vector-borne diseases. Meningitis, influenza, viral hepatitis, malaria, dysentery, and dengue have particular military significance because of their high epidemic potential and associated morbidity.

c. Other diseases of military concern have less explosive impact, but they do have high public health importance due to transmissibility. These include sexually-transmitted diseases (STDs), human immunodeficiency virus (HIV) infection, and tuberculosis. Other diseases, such as rabies and viral hemorrhagic fever, may become particularly significant in epidemic situations due to the high mortality associated with such infections.

d. Preventive measures include personal protective measures (for example, personal hygiene, immunizations, prophylactic medications, and repellents) and environmental control measures (for example, disinfection of water supplies, proper food handling practices, area vector control, and other aspects of field sanitation). Effective implementation of preventive measures require command emphasis and command, unit, and individual soldier education on ways to prevent illnesses.

4–2. Guidance
a. Disease prevention and control measures will ordinarily be determined at DA level and be set as the standards for the Army. Unless otherwise directed by HQDA, the principles and procedures recommended in the prevention and control of communicable diseases, as specified in the latest edition of Control of Communicable Diseases in Man, published by the American Public Health Association (field manual (FM) 833/NAVMED P5038), will be followed. More specific guidance may be obtained from OTSG, ATTN:SGPSPSPD.

b. PVNTMED services and teams will be familiar with disease prevention and control measures and will provide advice and guidance to commanders, units, and individuals on the prevention of communicable diseases. PVNTMED services and teams will also provide guidance to units on disease and environmental threats, specific preventive measures, and medical surveillance during and following deployments.

4–3. Functions
a. All commanders will—
   (1) Direct the institution and implementation of required disease preventive and control measures.
   (2) Ensure compliance of all eligible personnel with prescribed individual protective measures.
   (3) Ensure compliance with immunization requirements under AR 60020 and AR 40562/NAVMEDCOMINST 6230.3/AFR 16113/CJ COMDTINST M6230.4D.
   (4) Provide protective clothing, equipment, supplies, and facilities when required.
   (5) Provide orientation to their units regarding the prevention of heat and cold injuries (app B).

b. IMAs will—
   (1) Identify potential disease and environmental threats and/or conditions of epidemic potential, based on epidemiological information, medical intelligence, and knowledge of military activities.
   (2) Recommend individual protective measures and environmental control measures to the command, based on the health threat.
   (3) Conduct continuous medical surveillance of individuals and units in high risk situations and/or units operating in environments where the potential for acquiring serious infectious diseases is significant.
   (4) Conduct epidemiological investigations of suspected disease outbreaks or disease occurrences capable of reducing military effectiveness or readiness.
   (5) Report unusual occurrences of diseases or environmental health problems to commanders so corrective action can be taken immediately.

   c. Individuals will—
      (1) Comply with all preventive measures defined by command authorities.
      (2) Avoid unnecessary exposure to infectious agents, hosts, or vectors of disease.
      (3) Practice good personal hygiene.

4–4. Immunization and chemoprophylaxis requirements
For policies and procedures to be followed in immunization and disease chemoprophylaxis, see AR 40562/NAVMEDCOMINST 6230.3/AFR 16113/CJ COMDTINST M6230.4D. Immunization requirements for Active Duty and Reserve personnel contained in AR 40562/NAVMEDCOMINST 6230.3/AFR 16113/CJ COMDTINST M6230.4D or as directed by HQDA take precedence over guidance provided by the U.S. Public Health Service (USPHS) or the Centers for Disease Control, Atlanta, Georgia.

4–5. Specific programs
a. **Acute respiratory disease (ARD).**
   (1) ARD can result in considerable manhours lost due to morbidity from various infectious agents and their high transmission potential. Agents of greatest military significance are: Influenza, parainfluenza, adenoviruses, streptococcal infections, and mycoplasma infections. Other viral and bacterial agents are capable of causing ARD.
(2) The ARD season in the northern hemisphere normally extends from October through March, although cases can occur throughout the year. Recruits and personnel living in confined quarters are particularly susceptible to outbreaks of ARD.

(3) The Army ARD Surveillance and Control Program will be well-defined at all basic training installations and will consist of—

(a) Monitoring of ARD rates among basic trainees on a daily and weekly basis. Reports will be provided to appropriate higher headquarters commands on a regular basis.

(b) Monitoring of Group A streptococcal infections among basic trainees. Particular attention will be directed at changes in throat culture recovery rates and the presence of rheumatogenic strains of Group A streptococcal organisms. High prevalence of streptococci may indicate the need for penicillin (bicillin) prophylaxis. Such prophylaxis will be instituted at the direction of HQDA.

(c) Influenza and adenovirus immunizations to all recruits (AR 40562/NAVMEDCOMINST M6230.3M). See also M6230.4D.

(d) Implementation of any new surveillance or immunization program directed by HQDA.

(4) All Army installations will implement a monitoring system to detect unusual outbreaks of ARD. This ordinarily will require coordination with clinics and emergency rooms and monitoring of overall rates of school or work absenteeism, particularly during the ARD season. IMAs will report any unusual ARD activities by Special Telegraphic Report of Selected Condition (RCS MED16).

(5) Commanders will monitor compliance with the annual influenza immunization program to ensure a high level of participation.

Meningococcal infection.

(1) Meningococcal infection is associated with significant morbidity and mortality with the development of life-threatening meningitis and/or disseminated infection. The disease is highly transmissible through the airborne route, and is, therefore, a disease of special concern in those environments where personnel are confined in limited space or are in close physical contact with each other, as in household settings.

(2) The recruit training environment is of particular concern and has been the site of many outbreaks in the past. Meningococcal vaccine is, therefore, routinely administered year-round to basic trainees. HQDA will define other settings where meningococcal vaccine should be routinely administered.

(3) IMAs will be familiar with the requirements for immunization and chemoprophylaxis specified in AR 40562/NAVMEDCOMINST M6230.3M/AFR 16113.CG COMDTINST M6230.4D.

(4) Meningococcal infections will be promptly reported by RCS MED16 (AR 40400). All Neisseria meningitidis specimens will be submitted for group typing. Specimens subcultures will be shipped to: The Walter Reed Army Institute of Research, Bacterial Disease Division (ATTN: SGRDUWFA), Washington, DC 203075100. Information on the patient’s status should be included. Prior to shipment, laboratories should call the Bacterial Disease Division at AUTOVON 2913303.

(c) Malaria.

(1) Malaria represents a major threat to military readiness. Malaria chemoprophylaxis will be instituted when personnel are at risk of contracting malaria. Although falciparum malaria is generally regarded as a more serious potentially life-threatening type of malaria, vivax malaria is also considered a military threat due to its ability to incapacitate susceptible hosts. Therefore, chemoprophylactic measures will be directed against both forms of malaria.

(2) Specific drugs to be used will be based on the prevalence of specific types of malaria in the area of anticipated travel and the current drug resistance patterns. These prophylactic drugs include chloroquine, primaquine, chloroquine-primaquine (combination form), doxycycline, Fansidar, and mefloquine (investigational drug status). Guidance on the most appropriate chemoprophylactic medication will be provided by HQDA (contact the Disease Control Consultant, AUTOVON 2890125).

(3) Commanders will—

(a) Ensure that all personnel in their command receive health education on the prevention of malaria, and are aware of the need to seek medical attention should they experience any febrile illnesses during or following assignment to malarious areas.

(b) Ensure that all individuals in their command are taking appropriate drug prophylaxis during and following periods of travel to malarious areas.

(c) Ensure that preventive measures, including the use of military-approved repellents, the use of bednetting, and the proper wearing of protective clothing, are followed. In some situations, mosquito control measures may be indicated; such area control measures will be conducted by appropriate preventive medicine teams.

(4) IMAs will report any suspected or confirmed cases of malaria by RCS MED16 (AR 40400).

(d) Viral hepatitis.

(1) Viral hepatitis may be caused by several different viruses capable of causing liver failure. Hepatitis A and non-A non-B hepatitis are transmitted by the fecal-oral route, while hepatitis B is transmitted through the exchange of blood or other body fluids from an infected person.

(2) The Army Hepatitis Prevention and Control Program consists of—

(a) Community and unit health education, incorporating principles of good personal hygiene and sanitation.

(b) Administration of immune globulin to personnel considered to be at risk of contracting hepatitis A (AR 40562/NAVMEDCOMINST M6230.3M/AFR 16113.CG COMDTINST M6230.4D).

(c) Immunization of all active duty AMEDD personnel and other people considered to be at risk of contracting hepatitis B (health care workers, spouses or sexual contacts of hepatitis B carriers, newborns of hepatitis B carrier mothers, and close contacts of persons known to be carrying the hepatitis B virus).

(d) Immunization of Army personnel on permanent change of station (PCS) moves to the Republic of Korea, where hepatitis B is highly prevalent.

(e) Immunization of other military personnel considered to be at risk of contracting hepatitis B infection, such as selected Special Forces personnel.

(f) Prenatal screening for the presence of hepatitis B surface antigen.

(g) Screening of donated blood for the presence of hepatitis B virus and other screening procedures recommended by the American Association of Blood Banks, and the removal of suspected contaminated units from the inventory.

(h) Medical evaluation and counseling of all suspected and confirmed cases of hepatitis, to include acutely ill individuals and chronically infected persons.

(3) IMAs will conduct an epidemiological investigation in all cases of viral hepatitis. Outbreaks of hepatitis will be reported by RCS MED16 (AR 40400).

(e) Sexually-transmitted diseases.

(1) STD are defined as those infections that can be transmitted through sexual contact and for which sexual transmission is epidemiologically important. Once categorically referred to as venereal diseases, STD include: Gonorrhea, syphilis, chancroid, chlamydia, genital herpes, lymphogranuloma venereum, granuloma inguinale, venereal warts (condyloma acuminate), and non-gonococcal urethritis. Hepatitis B and HIV infections are also considered to be STD.

(2) The Army STD Prevention and Control Program consists of—

(a) Accurate diagnosis and appropriate treatment of infected persons and their sexual partners (AR 600110 and AR 600240/BUPERINST 1752.1/AFR 21118/MCO 1752.1).

(b) Personal interviews and epidemiological contact investigation.

(c) Active surveillance at the installation level.

(d) Health education directed at all sectors of the military community.

(e) Control of prostitution (AR 21010).

(3) At the installation level, STD prevention and control efforts
include appropriate therapy and follow-up, disease intervention, identification of locations where a high level of STD transmission may be occurring, and community and unit health education. Centralization of diagnostic efforts, interviewing and counseling, and treatment procedures is ideal and lends itself to better quality control and maintenance of patient confidentiality.

(4) Unit health education classes are strongly encouraged and should be incorporated with HIV education efforts and classes on personal hygiene whenever possible.

(5) Punitive action against a person will not be based solely on the fact that an individual has contracted an STD. However, in selected situations, repeated occurrences of STD may be the basis for administrative and/or corrective action under appropriate Army regulations.

(6) IMAs will collect on a monthly basis STD statistics and report these statistics to higher headquarters. STD statistics should not be unit-specific but will include the major categories of STD (rate/1000/month).

(7) STD information and/or statistics will not be used to compile indices of unit morale or integrity or commander efficiency.

(8) The release of medical information concerning persons who have contracted an STD will be based on applicable law and regulations. The Seven Point Agreement of 1967 (An Agreement on Measures for the Control of Venereal Diseases) between the Department of Health and Human Services (formerly Department of Health, Education, and Welfare), Department of Transportation, the Association of State and Territorial Health Officers, and the DOD is in effect. Also involved are cooperation with civilian health authorities and the reporting of military statistics to State or local health departments.

(9) The success of the military program is contingent on a satisfactory working relationship with civilian public health authorities. A cooperative atmosphere with local, county, and State health officials involved in the prevention and control of STD in an area is encouraged.

(10) HQDA will provide periodic guidance on the recommended treatment for uncomplicated gonorrhea and other STD. These guidelines take precedence over USPHS guidelines.

f. Rabies.

(1) Rabies is almost 100 percent fatal. Although the incidence is low, the possibility of animal bites in military personnel is real. Each bite incident must be carefully evaluated by medical authorities involved in rabies prevention and control efforts.

(2) The Army Rabies Prevention and Control Program consists of—

(a) Rabies pre-exposure prophylaxis with human diploid cell rabies vaccine. This prophylaxis will be administered to persons considered to be at significant risk of being bitten by potentially rabid animals(AR 40562/NAVMEDCOMINST 6230.3/AFR 16113/CJ COMDTINST M6230.4D).Veterinarians, animal trappers, and selected animal control officers are included in this category of personnel at occupational risk of exposure. Selected military personnel in remote rabies-endemic areas should be considered for pre-exposure vaccination.

(b) Rabies post-exposure prophylaxis. Prophylaxis with human diploid cell rabies vaccine and human rabies immune globulin will be based on the incidence of animal rabies in the geographical area; the species of animal involved; the vaccination status of the animal; and the circumstances surrounding the bite incident.

(c) Stray animal control efforts.

(d) Continued surveillance of animal rabies in the area (domestic and wild animals in the geographical area).

(e) Community health education on the threat of rabies (domestic and wild animals and pets).

(f) IMAs will designate at least one qualified physician (usually the PVNTMED medical officer) and one veterinarian as rabies advisers. Rabies advisers will be consulted whenever the attending physician contemplates administration of specific anti-rabies treatment.

(4) Animal bite incident reports will be generated on every domestic and wild animal bite or suspected rabies exposure. These forms will be reviewed on a daily basis by one of the rabies advisers.

h. Tuberculosis.

(1) Tuberculosis screening programs detect previously unrecognized cases of active tuberculosis or persons exposed to tuberculosis who may have inactive infection. Since as many as 1 in 10 infected persons may develop the disease at some time in their lives, and risk is greatest in the first year after infection, the Army Tuberculosis Surveillance and Control Program has been established to identify persons who have been infected with the tubercle bacilli.

(2) Screening is based on the use of the tuberculin skin test(TST). Types of TSTs and directions on the administration of these tests are available through the American Thoracic Society or the USPHS. The intradermal Mantoux test is the standard test and should be read 48 to 72 hours after application.

(3) Appropriate drug treatment can be given to infected persons identified through screening, and prophylaxis against the development of active disease can be administered to those at risk. Isoniazid (INH)administered orally is normally used for preventive therapy (300 mg daily for adults and 10 to 14 mg/kg body weight not to exceed 300 mg daily for children). Other prophylactic drugs, such as rifampin, should only be used in those selected situations when there is a well-defined high risk exposure to a patient with infectious tuberculosis who is excreting INH-resistant organisms.

(4) For personnel not previously known to have a positive TST, skin tests will be administered to—

(a) Personnel entering active duty for 30 days or more as part of reception processing.

(b) Military and civilian personnel travelling OCONUS on PCS orders under DA auspices. TST will be applied within 3 months of anticipated move OCONUS.

(c) Military and civilian personnel returning to CONUS from an OCONUS assignment within 2 months of return. If an individual fails to receive a TST prior to departure or return, one will be administered within 2 weeks of arrival at the next duty station.

(d) Military and civilian personnel undergoing periodic physical examinations (ordinarily every 5 years for most military members), unless one has been administered within the past 6 months.

(e) Prospective civilian employees, as a condition for employment in health care facilities, schools, child development services (CDS), or other environments where tuberculosis may constitute a special hazard to others.

(f) Health care or community service personnel who may be at increased risk of contracting tuberculosis, periodicity of testing will be determined locally by IMAs, based on risk of exposure in specific occupational settings.

(g) Healthcare beneficiaries undergoing medical evaluation for tuberculosis, HIV infection, or other diseases associated with tuberculosis.

(5) TST reactors are defined as individuals who have a positive skin test of 10 mm induration or greater to the intermediate strength Mantoux test. TST converters are reactors whose TST has changed from negative or doubtful to positive (greater than 10 mm induration) within the past 2 years.

(6) For individuals known to have a positive TST previously, no further TSTs need be applied. Medical records will reflect the positive TST status, and an annotation as to the medical evaluation performed recently or in the past will be made. If INH therapy had been instituted, this information, along with details of duration of therapy and recommendations for follow-up, will be recorded on the records.

(7) For individuals identified for the first time as TST-positive
with no clear history of when conversion occurred, a medical evaluation will be performed to determine if active disease is present. The evaluation will include a careful medical history eliciting signs or symptoms suggestive of infection and a chest x-ray (alveolar, lateral, and apical lordotic views). This information will be entered in the medical record.

(8) All individuals with a recently identified positive TST must be evaluated and considered for INH chemoprophylaxis. Persons under age 35 should be placed on INH prophylaxis routinely, unless medical conditions exist that contraindicate its use. Individuals 35 years and older should be carefully evaluated, but, because of a higher association of INH-induced hepatitis in older individuals, the decision to begin INH prophylaxis should be made on a case-by-case basis based on risk and time of likely conversion.

(9) Duration of INH chemoprophylaxis for TST reactors with no risk factors for the development of active disease may be shortened from 12 months to 6 months. This includes persons with TST reactions of 5 mm or greater induration and who are close household contacts of active cases. Contacts of infectious cases with negative TST reactions of less than 5 mm induration should continue daily INH for 3 months; if the TST remains negative, prophylaxis can be discontinued.

(10) TST reactors with risk factors for the development of active tuberculosis should be placed on INH for 12 months. Risk factors include—

(a) Abnormal chest x-ray with stable parenchymal lesions.
(b) HIV infection.
(c) Adrenocorticosteroid or immunosuppressive therapy.
(d) Reticuloendothelial/hematologic disorders such as leukemia, lymphoma, or sickle cell anemia.
(e) Diabetes mellitus.
(f) Sarcoidosis.
(g) Chronic hemodialysis.
(h) History of gastrectomy.
(i) Chronic undernutrition or weight loss.

(11) Liver transaminase tests will be performed at the beginning and periodically during therapy; that is, at 1 and 3 months. If transaminase levels exceed 3 to 5 times the upper limit of the normal range of the laboratory, the decision to continue INH prophylaxis should be reconsidered.

(12) TST sensitivity and immunity to tuberculosis after Bacille Calmette-Guerin (BCG) vaccine is highly variable, and there is no reliable method for distinguishing tuberculin reactions caused by BCG from those caused by natural infections. Since the incidence of tuberculosis is high in countries with BCG vaccination programs, a positive TST should be evaluated independently of BCG history.

(13) TST results will be documented on the SF 601 (Health Record) and on HEW Form PHS 731 (International Certificates of Vaccination).

(14) A local tuberculosis registry of all persons under medical surveillance will be maintained. This registry will serve as a current listing of all active and inactive cases and contacts requiring medical follow-up. DA Form 3897R (Tuberculosis Registry) will be used for this purpose and will be locally reproduced. DA Form 3897R is located at the back of this regulation.

(15) For personnel under surveillance undergoing a change of assignment, DA Form 3897R will be mailed to the medical commander of the gaining organization to ensure continuity of care. The individual will be counseled prior to his or her departure.

(16) For personnel under surveillance departing military service, IMAs will notify the appropriate State health department where the individual will be living. The Veterans Administration ordinarily assumes responsibility for military separatees who are under active tuberculosis surveillance.

(17) When medical surveillance is discontinued, DA Form 3897R will be placed in a closed file, retained for 1 year, then destroyed per AR 254002.

Section II
Climatic Injury Prevention and Control

4–6. General

a. Environmentally-associated illnesses may be present as isolated events or be of epidemic proportions with significant military impact. Included are heat injuries, cold injuries, and high altitude sickness. Other environmental conditions based on specific scenarios may justify the need for additional interventions such as vision protection against excessive glare or skin, eye, and respiratory tract protection from wind or excessive dust or sand.

b. The goal of this program is to prevent injuries from ever occurring though a comprehensive program incorporating health education, personal protection, and appropriate adjustment of activities as indicated, to include acclimatization in hot environments. Monitoring of the environment is critical. Commanders, through early recognition of climatic injuries and prompt implementation of additional preventive measures, can reduce morbidity in the command.

4–7. Functions

a. Commanders will—

   (1) Provide protective clothing, equipment, supplies, and facilities (other than medical) to prevent climatic injuries from occurring.
   (2) Implement heat and cold injury prevention programs within their respective commands, the actions being based on guidelines described in paragraph 46.
   (3) Monitor environmental conditions where troops will be located with assistance provided by medical authorities.

b. IMAs will recommend individual or environmental protective measures to the command.

Section III
Hospital Infection Control

4–8. General

An effective program for the prevention and control of hospital-associated infections is required to meet the objectives of high quality patient care and the effective utilization of hospital services. This program encompasses virtually all hospital operations under the responsibility of the hospital commander. The hospital commander implements these objectives. The hospital infection control committee supervises the program.

4–9. Hospital infection control committee

a. Committee policy and responsibility. The committee operates as a medical audit committee and, as part of the quality assurance program, is responsible to the commander through the QA committee. The basic principles of patient care and employee health inherent in the program apply to all inpatient areas, outpatient areas, emergency rooms, special care areas, and troop clinics.

b. Objectives. The committee’s primary objectives are as follows:

   (1) Ensuring a continuing education program on the control of hospital-acquired infections.
   (2) Establishing a practical and timely system for the recognition, evaluation, and reporting of nosocomial infections in hospitalized patients and recently discharged patients.
   (3) Providing assistance in developing preventive measures and policies.
   (4) Reducing the incidence of preventable infections.

c. Composition. The committee chairman will be a medical officer who demonstrates interest and knowledge in infection control, and can devote sufficient time and effort to ensure an effective program. Committee membership should include the following:

   (1) Hospital epidemiologist. Each hospital commander will designate an officer with appropriate training and experience as the hospital epidemiologist. This officer will supervise the educational program and surveillance activities, assist in monitoring infection control policies and procedures, and serve as adviser to the hospital commander, medical staff, and nursing staff on infection control practices.
(2) **Hospital infection control officer.**
   (a) This officer, appointed by the hospital commander and normally a nurse, will serve as liaison between the hospital infection control committee and all departments or services of the hospital to—
   1. Facilitate clinical and environmental surveillance activities.
   2. Foster an attitude of cooperation.
   3. Enhance the effectiveness of the educational program.
   (b) At the direction of the chairman, hospital infection control committee, the hospital infection control officer will coordinate all educational activities, and gather clinical data to determine the incidence of endemic infections (para 411) and manage epidemic events. This includes their epidemiological investigation and reporting.
   (c) The hospital infection control officer can also serve as the hospital epidemiologist.

(3) **Clinical service representatives.** Representatives of the major clinical departments and services, including nursing personnel, may serve as members of the hospital infection control committee to provide the necessary interdisciplinary clinical input. Representation from the house staff, when applicable, is desirable and encouraged.

(4) **Environmental science officer (AOC 68N).** The environmental science officer is normally the principal adviser on matters relating to the hospital environment including waste management, ventilation, housekeeping, selection, and use of antiseptics and disinfectants, food sanitation, linen management, and environmental monitoring.

(5) **Administrative officer.** The hospital executive officer is normally the principal adviser on administrative matters and services.

(6) **Microbiologist.** The microbiology/serology section of the clinical laboratory should be represented to provide the necessary input on microbiological data and procedures.

(7) **Entomologist (AOC 68G).** When available, the entomologist will be the principal adviser on the potential for pest infestations that contribute to the spread of infectious agents and will also advise on the implementation of proper pest control measures. When an entomologist is not available, the environmental science officer will function in this role.

(8) **Other consultants.** Representatives of the pharmacy, food, and housekeeping services, as well as other consultants, should be available for infection control committee meetings when required.

d. **Functions.** Committee meetings will be held at least every 2 months or as often as necessary to accomplish the objectives. The committee will—
   (1) Describe standard criteria for defining nosocomial infections.
   (2) Establish written policies and procedures relating to isolation techniques, antiseptics, disinfection and sterilization techniques, waste management (para 116), and general sanitation.
   (3) Establish written policies and procedures concerning patient care techniques and measures for the prevention of infections in patients and personnel.
   (4) Ensure that policies and procedures developed for such activities as clinics, special care services, laboratories, and support services adequately address the potential for infections and their prevention.
   (5) Provide for a review, at least annually, of all hospital and clinic written policies and procedures related to infection control and to determine their applicability and to revise as appropriate.
   (6) Provide assistance in the development of the infectious disease aspects of the hospital employee health program. (See chap 5.)
   (7) Coordinate with the medical staff in its review of the clinical use of antimicrobial agents by analyzing and using significant surveillance data and antimicrobial susceptibility test data.
   (8) Recommend to the hospital commander actions that should be taken to control hospital outbreaks of infectious diseases.

d. **Education.** Provisions will be made by the hospital infection control committee for the orientation of all new hospital personnel to their responsibilities in the prevention and control of hospital-associated infections. Periodic inservice education in infection control will be provided in all departments and services and will be documented. Information, including data supporting significant trends, will be reported to the MEDDAC or MEDCEN QA committee and be incorporated into departmental educational programs as well as in formal presentations of the most current prevention and control concepts to the medical staff (see AR 4068).

4–10. **Technical assistance**

On-site consultations and special studies should be requested from the hospital epidemiologist or hospital infection control officer at the relevant MEDCEN; for example, Fort Carson would contact the appropriate individuals at Fitzsimons Army Medical Center. The normal consultation routes will be as follows: MEDDAC to regional MEDCEN; regional MEDCEN to either TSG’s physician or nurse consultant for hospital infection control. However, requests may also be made directly to OTSG.

4–11. **Reporting**

a. An endemic nosocomial infection rate for the hospital will be consolidated into formal reports for presentation during medical staff conferences and for inclusion in CHRs. Nosocomial infections for a suitable period of time will be reported in CHRs at least three times a year by the total surveillance (incidence) or prevalence rate as follows:
   (1) The total surveillance (incidence) rate equals—
      (a) The number of nosocomial infections per unit of time.
      (b) The number of patients discharged per unit of time.
   (2) The point prevalence rate equals—
      (a) The number of patients with nosocomial infection at the time of the survey.
      (b) The number of patients in the hospital at the time of the survey.
   (3) The number of patients with multiple nosocomial infections will be listed.

b. Coding of diagnoses on individual patient data system coding transcripts from inpatient treatment record cover sheets will always include any diagnosis representing a hospital infection (AR 4066).

c. Certain infections of high communicability as well as significant outbreaks of infection will be reported expeditiously by RCS MED16 (chap 3).

**Chapter 5**

**Occupational Health Program**

**Section I**

**General**

5–1. **Background**

This chapter prescribes the Occupational Health Program and services required under provisions of Executive Order 12196 and DOD Instructions 6050.5, 6055.1, 6055.5, and 6055.12 for DA military and civilian personnel.

5–2. **Objectives**

The objectives of the Army Occupational Health Program are to—

a. Assure that all eligible personnel (military and civilian) are physically, mentally, and psychologically suited to their work at the time of their assignment, and that physical and mental health are monitored to detect early deviations from the optimum.

b. Protect employees against adverse effects of health and safety hazards in the work environment. This includes field operations as well as the industrial workplace.

c. Assure proper medical care and rehabilitation of the occupationally ill and injured.

d. Reduce economic loss caused by physical deficiency, sickness, and injury of civilian employees.

e. Prevent decreased combat readiness caused by occupational illness and injury of military personnel.
5–3. Army Occupational Health Program

a. The overall Occupational Health Program promotes health and reduces risk of illness arising from the individual’s work environment. This encompasses special preventive measures for both military and civilian personnel who are exposed or potentially exposed to toxic materials, infectious agents, or other hazardous influences of the work environment.

b. Medical measures will be carried out according to professional standards in the field of OH.

c. Army occupational safety and health standards are noted in (1) through (5) below. When alternate or supplemental standards are necessary, documentation with justification will be forwarded through command channels to HQDA (SGPSPSP), 5109 Leesburg Pike, Falls Church, VA 220413258, to obtain appropriate approval authority.

(1) DOD and DA OSH standards for military and nonmilitary workplaces for which regulatory agencies either have or have not issued OSH standards. This includes DOD and DA pamphlets, circulars, technical bulletins, and messages.

(2) OSHA standards including emergency temporary standards with minor adaptation as necessary to conform with DA administrative practices.

(3) Alternate workplace standards based on publications relating to workplace exposure criteria. These standards may be used in lieu of existing OSHA standards or in which no OSHA standard exists. The current American Conference of Governmental Industrial Hygienists threshold limit values will be the standards used in DA military and civilian workplaces if—

(a) OSHA standards are less stringent.

(b) No OSHA standard exists.

(4) Other regulatory workplace standards issued under statutory authority by other Federal agencies (such as the Department of Transportation and the Environmental Protection Agency (EPA)).

(5) Special DA OSH standards developed for military-unique equipment, systems, and operations.

A viable Occupational Health Program requires continuing cooperation among managers, supervisors, personnel officers, and safety and medical personnel to include division surgeons, optometrists, industrial hygienists, audiologists, and safety personnel.

e. As a minimum, the Occupational Health Program will include the following elements:

(1) Inventory of chemical, biological, and physical hazards in the work environment of all installation activities, including MTFs and research and development activities.

(2) Job-related medical surveillance.

(3) Administrative medical examination.

(4) Employee education about job-related health hazards.

(5) Treatment of occupational illness and injury and emergency treatment of nonoccupational illness and injury.

(6) Hearing conservation.

(7) Occupational vision.

(8) Pregnancy surveillance.

(9) Job-related immunizations.

(10) Illness absence monitoring.

(11) Chronic disease surveillance.

(12) Epidemiologic investigations of occupational illness and injury.

(13) Maintenance of OH medical and administrative records and reports.

(14) Industrial hygiene surveys and safety and health inspections.

f. Other services that may be provided when adequate resources are available include but are not limited to—

(1) Group counseling on specific problems or habits affecting health.

(2) Disease screening.

(3) Voluntary periodic health examinations on an age-related basis.

5–4. Program functions

a. The installation commander will ensure that—

(1) Employees under his or her command are provided OH services required by this chapter.

(2) Supervisors at all levels are informed of and carry out their responsibilities in the program.

(3) Individual employees are informed of potential OH hazards and safe practices and procedures, and are instructed in the wearing of PPE.

(4) An Occupational Health Program administrator or coordinator will be designated at installations or activities that do not have an occupational health clinic.

(5) A program for the recognition, evaluation, and control of unhealthful working conditions is established. This program will include—

(a) Publishing a local regulation or supplement to an existing regulation that delineates the responsibilities of all installation OSH participants.

(b) Ensuring establishment of a safety and occupational health (SOH) advisory council (AR 38510).

(6) The installation Asbestos Management Program is established per TB MED 513 and other DA guidance.

b. Safety manager responsibilities are defined in AR 38510 and AR 38540.

c. Civilian and military personnel officers will provide support and guidance to ensure efficient accomplishment of the overall program (AR 6008 and Federal Personnel Manual (FPM) chaps 250, 290, 293, 294, 339,792, and 810). This includes coordination with OSH personnel to ensure that—

(1) A suspense system is maintained to—

(a) Identify personnel in positions requiring specific standards of physical fitness and job-related medical surveillance.

(b) Schedule personnel for the indicated preplacement, change of position, periodic, fitness for duty, and termination examinations.

(2) Applicants and employees are advised regarding potential OH hazards, appropriate protective equipment, safety practices, and job-related medical surveillance requirements of their work assignments.

(3) The commander of the MTF providing medical support will—

(1) Program resources to ensure provision of OH services required by this regulation.

(2) Ensure provision of physician support for OH services where there is no physician assigned.

(3) Appoint an audiologist, when available, to act as the hearing conservation officer and to participate as a member of the SOH advisory council. If an audiologist is unavailable, the IMA will designate an individual from the occupational medicine staff to act as the hearing conservation officer. (See the definition of occupational medicine staff in the glossary.)

(4) Appoint an individual to act as the industrial hygiene program manager according to TB MED 503.

(5) Project the impact of full-scale industrial mobilization on OH services and ensure provision of these services through the use of contingency contracts and on expanded mobilization TDA.

e. The chief, PVNMTMED service, will—

(1) Provide overall technical guidance for the Occupational Health Program to appropriate supporting clinical services and to the tactical unit surgeons.

(2) Ensure proper coordination with installation and MTF safety and personnel offices, hospital infection control personnel, and the division surgeon.

(3) Initiate, if appropriate, and assist in epidemiologic investigations.

(4) Ensure maximum use of the military occupational health vehicle, where applicable, to conduct monitoring audiometry.

(5) Support the installation Asbestos Management Program according to TB MED 513.

(6) Provide medical review of Federal Employees Compensation Act claims.

f. The SOH advisory council committee (as described in AR 38510) will—

(1) Consider matters involving OSH.

(2) Make recommendations to the installation commander.

(3) Perform such additional tasks as the commander or council
chairperson may direct. DOD components may exempt installations with a very small population from the requirement to establish a council.

(4) Review, discuss, and make comments on the installation OSH hazard abatement plan or schedule.

g. The OH representative of the SOH advisory council committee will provide input concerning specific health aspects of council responsibilities. The representative will—

(1) Provide information and make recommendations concerning required actions to implement applicable laws and regulations related to health.

(2) Provide advice, guidance, and/or coordination on required actions to comply with survey and inspection recommendations made by higher headquarters and other agencies.

(3) Provide the council with data regarding accident and illness trends and bring to the council’s attention any problems related to employee participation in job-related health programs.

h. The OH nurse and the CHN, as deemed necessary by the chief of the PVNTMED service, will coordinate Occupational Health Program activities in the areas of epidemiology, educational programs, communicable disease programs, and use of community resources. The activities will include but are not limited to the following:

(1) Reviewing DA Form 3076 to detect illness and injury patterns.

(2) Providing advice, as needed, in matters pertaining to OH needs of soldiers.

i. The chief, OH, will—

(1) Plan, direct, supervise, and evaluate the Occupational Health Program according to specific installation needs and resources and requirements of this regulation.

(2) Coordinate with other MTF and installation staff (including labor relations advisers) and with the division surgeon to ensure—

(a) Provision of required OH services.

(b) Collection, review, and reporting of required OH data.

(3) Conduct or coordinate medical surveillance and health hazard training for military and civilian employees potentially exposed to OH hazards, and evaluate employees in positions requiring specific standards of physical fitness.

(4) Regularly visit work areas to keep informed about work operations and potential hazards and maintain working relationships with supervisors and employees.

(5) Conduct epidemiologic investigations of actual or suspected occupational illness.

(6) Provide advice and guidance to commanders and other concerned personnel (such as employee representatives) regarding OH matters.

(7) Participate in the installation SOH advisory council committee and quality control committee.

(8) Establish a light duty or limited duty program, in coordination with the installation commander, safety officer, and personnel officer, to facilitate an early return to work for employees injured on the job.

j. The industrial hygienist will—

(1) Develop and update annually industrial hygiene input into the Occupational Health Program document to clearly define goals and objectives in the industrial hygiene area.

(2) Establish and maintain the HHIM of the OHMIS.

(3) Develop an industrial hygiene implementation plan for the allocation and application of industrial hygiene resources.

(4) Perform industrial hygiene evaluations of workplaces, provide technical guidance and support for the hazard communication, asbestos abatement, and installation OH programs, and perform other responsibilities as defined in TB MED 503.

k. The chief, DPCCM, will provide clinical support and coordinate with the chief, PVNTMED, and the chief, OH, to assure provision and reporting of required OH services for military and civilian employees.

l. The chief, optometry, MEDDAC, will serve or appoint an optometrist as the occupational vision officer who will—

(1) Assist OSH personnel in identifying eye-hazardous occupations, areas, tasks, or processes and in determining the type of protective eyewear required.

(2) Ensure that verification of prescription and proper fitting of industrial safety spectacles are accomplished.

(3) Ensure that industrial safety spectacles meet current American National Standards Institute (ANSI) Z87.1 criterion.

(4) Assist OH personnel in establishing and maintaining a vision screening program for workers in potentially eye-hazardous occupations and other vision screening programs when required.

(5) Provide professional vision evaluations and the necessary spectacle corrections for civilian employees referred under the Occupational Vision Program.

(6) Provide technical input and assistance for the Employee Health Hazard Education Program.

(7) Provide professional guidance regarding the wearing of contact lenses in the industrial environment. Contact lenses provide very limited industrial eye protection; therefore, proper protective eyewear should be used.

m. The chief, patient administration division, will act as technical adviser for patient administration aspects of the Occupational Health Program, to include collection and use of required OH reports data.

n. Managers and supervisors at all levels will—

(1) Keep informed about OH hazards and the medical and safety requirements in activities under their control.

(2) Train and educate employees regarding job health hazards and appropriate safety practices.

(3) Enforce the use of protective clothing and equipment.

(4) Advise the IMA of proposed or actual changes in work operations that may affect the health or safety of the worker.

(5) Provide the civilian and military personnel offices with the health and safety information necessary for effective job classification and placement actions.

(6) Assure that employees are referred for required job-related medical surveillance.

(7) Employees will—

(1) Follow safe and healthful work practices.

(2) Use PPE when required.

(3) Make note of and report suspected unsafe or hazardous work situations.

(4) Comply with requirements of the Occupational Health Program.

Section II

Occupational Health Management Information System

5–5. General

a. The purpose for OHMIS is to assist OH professionals in improving the effectiveness and economy of OH delivery through provision of accessible, timely, accurate data on both military and civilian employees, their workplace environment, and their health status. OHMIS is configured as a distributed network with installation level processing on microcomputers at each Army OH facility worldwide.

b. Three modules support the Army OH team:

(1) The Hearing Evaluation Automated Registry System will—

(a) Automatically perform hearing testing to include automatic forms completion; reduce lost training and work time.

(b) Significantly reduce error rates.

(c) Allow hearing conservation officers to quickly and easily determine program participation and hearing loss incidence and prevalence.

(2) The HHIM will—

(a) Maintain workplace descriptions including workplace hazard inventories, employee exposures, engineering and personal protective controls, and exposure abatement efforts.

(b) Document individual exposure histories.

(3) The Medical Information Module (MIM) will—

(a) Automate access to present and past exposure information, both workplace and individual employee.
(b) Automate access, verification, and update of demographic and clinical encounter information.
(c) Generate exposure-based recommended health surveillance procedures and provide locally tailored appointment schedules.
(3) All three modules combine to facilitate installation-level data base management, quality assurance, hazard communication and health education, resource management, and query and report preparation.

5–6. Functions
a. HSC is the designated Assigned Responsible Agency for operations, maintenance, and support of OHMIS after system deployment.
b. USAEHA is designated Proponent Agency for OHMIS. The OHMIS Coordinating Office has been established under the Director of Occupational and Environmental Health.
   (1) The functional program manager will—
      (a) Coordinate the separate module manager’s efforts for the maintenance and update of reference files that reflect Army policy regarding OH surveillance, exposure monitoring, and exposure definitions.
      (b) Evaluate the effectiveness of corporate and local OH programs and the conduct of corporate and local OH programs against established objective, discrete, measurable, and attainable performance standards adjusted for existing resources.
      (c) Respond, as appropriate, to queries for information derivable for the OHMIS database.
   (2) The individual module managers will respond directly to queries from the field relating to use or function of the three modules.
c. The Fort Detrick Director of Information Management is designated as the Application System Developer for OHMIS. As such, the Director will provide technical systems administration. The information center will provide assistance to users with hardware and/or software.
   d. At the installation level, the chief, PVNTMED services, will ensure systems administration security and the proper use of OHMIS.

5–7. Forms
a. The following occupational health patient forms are used to document both the workload requirements and activities of OH programs and the specific recipients of these activities:
   (1) Occupational Health Patient Form, DA Form 5931.
   (2) USEAER Occupational Health Form, DA Form 5932.
   (3) Occupational Health Patient Form—Supplemental, DA Form 5933 to be used in Panama, Puerto Rico, Guam, and the Virgin Islands, and Japan.
   (4) Korea Occupational Health Encounter Form, DA Form 5934.
   b. The data can be used to—
      (1) Obtain the information required for summary or statistical reports.
      (2) Help standardize Occupational Health Program elements and the services provided.
      (3) Initiate epidemiologic studies.
      (4) Increase program management efficiency.
      (5) Serve as a tracer or audit trail for services.
c. Instructions for preparing these forms are found in MIM’s Supplemental User’s Instructions, available from the Commander, USAEHA, ATTN:HSHBMOF, Aberdeen Proving Ground, MD 210105422.

Section III
Occupational Health Services

5–9. Medical examinations
a. Job-related examinations. Preplacement, job transfer, periodic, and termination examinations will be provided to all military personnel and civilian employees potentially exposed to health hazards in the work environment. Termination examinations will be provided on termination of assignment or termination of employment for all employees who have been included in a periodic job-related medical surveillance program unless an examination has been conducted within the past 90 days. The 90-day exception does not apply in cases where the content of the periodic examination differs from the termination examination, for example, high risk microwave or laser workers, or where a more stringent requirement exists. The chief, OH, or his or her representative, will review the HHIM (chap 5, sec II) annually and when operations change. Such a review is performed to determine the scope and frequency of job-related examinations for military personnel and civilian employees potentially exposed to health hazards. The medical surveillance matrix of the MIM will assist the OH care provider in making this determination by providing regulatory and recommended guidance for job-related examinations on each hazard in the matrix. The medical surveillance matrix is based on the hazards identified in the HHIM. The following documents will provide supplemental information:
   (2) National Institute for Occupational Safety and Health (NIOSH)Publication No. 81123.
   (3) DOD 6055.5M.
   (4) TB MED 501.
   (5) TB MED 502.
   (6) TB MED 506.
   (7) TB MED 509.
   (8) TB MED 510.
   (9) TB MED 513.
   (10) TB MED 523.
   (11) TB MED 524.
   (12) TB MEDs, DA pamphlets, and other documents concerning job-related medical surveillance requirements as they are developed and formally issued.

b. Military. In addition to routine entrance and periodic examinations performed under AR 40501, certain assignments will require further preassignment, periodic, and termination examinations that are specific for any potential chemical, physical, or biological hazards.
c. Civilian. In addition to job-related examinations required by a above, civilian employees assigned to positions requiring specific physical fitness standards will be provided examinations according to Office of Personnel Management (OPM) policy (FPM chap 339 and FPM chap 930). If necessary, job-related medical evaluations may be made a condition of employment. Employees not required to have preplacement examinations (FPM chap 339) should be scheduled for baseline health screening evaluations if resources permit. The baseline evaluations may include a health history, blood pressure determination, vision screening, and hearing tests.
   d. Other required examinations. Fitness for duty and disability retirement examinations will be accomplished according to FPM chapter 339. Medical examinations for individuals potentially exposed to chemical surety materials will be accomplished per the applicable DA pamphlets.
   e. Health maintenance examinations. While not a requirement for civilian employees, health maintenance examinations are encouraged, subject to availability of health resources. Such examinations may include single or multiple disease screening or more detailed medical evaluations, and can be offered on an age-related basis or to specific target groups.
   f. Follow-up. A follow-up system should be developed and maintained for all health examination and screening programs to identify and report their effectiveness and to assure indicated counseling and referral.

5–10. Illness and injury
a. Treatment for civilian employees.
1. Occupational illness and injury. Diagnosis and treatment of injury or illness sustained in performance of official duties is authorized by AR 40–3 and under the Office of Workers’ Compensation Program (FPM chap 810). Employees who request examination and treatment will be provided it at no cost at any Army MTF, other Federal MTF, or by a physician or hospital of his or her choice. If an Army dispensary, clinic, hospital, emergency room, or local facility under contract with the Army is available at the activity, locally prescribed procedures will require that the injured employee be initially referred to that MTF.

2. Nonoccupational illness and injury. Definitive diagnosis and treatment of nonoccupational illness and injury cases are not responsibilities of the Occupational Health Program except—

(a) In an emergency. The employee will be given the attention required to prevent loss of life or limb or relieve suffering until placed under the care of the employee’s personal physician.

(b) For minor disorders. First aid, or palliative treatment may be given if the condition is one for which the employee would not reasonably be expected to seek attention from a personal physician, or to reduce absenteeism by enabling the employee to complete the current work shift before consulting a personal physician. Requests for repetitive treatment of nonoccupational disorders will be discouraged.

(c) Minor treatments or services such as administering allergy treatments, monitoring blood pressure, providing physiotherapy, and so forth. These may be furnished at the discretion of the responsible physician if resources are available. A request must be submitted in writing by the employee’s personal physician. Medications, if required, must be provided by the employee.

(d) In cases of employees with an alcohol or drug abuse problem. These employees should be encouraged to seek assistance and counsel from the alcohol and drug abuse prevention and control program. AR 600–85, FPM Supplement 792–2, and DA Pam 600–17 provide further guidance. OH functions in this program include initial counseling and referral of employees to treatment resources.

b. Treatment for military personnel. AR 40–3 authorizes diagnosis and treatment for both occupational and nonoccupational illness and injury for military personnel. All incidents of military noncombat job-related illnesses and injuries will be reported to the proper occupational health and safety officials. Definitions of reportable occupational illnesses and injuries are found in AR 385–40, chapters 2 and 4, and in that publication’s glossary.

c. Medical directives. Comprehensive medical directives for emergency care and treatment of occupational and nonoccupational illnesses and injuries by the nursing staff will be prepared, signed, annually reviewed, and revised (if necessary) by the responsible physician to—

(1) Assure proper handling of emergencies in the absence of, or prior to, the arrival of a physician.

(2) Direct the care to be given for minor incidents not requiring personal attention of a physician.

(3) Authorize other activities by the nursing staff.

(d) First aid. In general, the placement of first-aid kits in work areas is discouraged. Exceptions should be made where work areas are geographically located distant from an MTF or where extremely hazardous exposures may occur and require immediate treatment for exposure. If first-aid kits are placed in work areas, their contents, intended use, and maintenance will be approved by medical personnel. Personnel rendering first-aid treatment will have approved first-aid training. All first-aid treatment rendered will be reported to OH personnel.

5–11. Epidemiologic investigations

Such investigations will be conducted after the occurrence of suspected or proven occupational illnesses. Identification of apparent excessive numbers of occupational injuries will be reported to safety officials. Investigations will be made, in coordination with safety officials when indicated, of employee reports of unhealthful working conditions. Situations that represent an imminent danger to Army personnel will be reported under AR 385–10.

5–12. Immunizations and chemoprophylaxis

a. Appropriate immunizations will be provided personnel with increased risk of infection related to potential job hazards or when required for official foreign travel. Other immunizations may be offered to civilian personnel to reduce absence due to sickness. Immunizations offered to civilian personnel will be based on current recommendations published by the USPHS (AR 40–562/NAVMED-COMINST 6230.3/AFR 161–13/CG COMDTINST M6230.4D).

b. Civilians traveling under military sponsorship will be provided appropriate immunizations and chemoprophylactic medications.

5–13. Illness absence monitoring

Medical support of the illness absence monitoring program for civilian employees will include—

a. Screening, treatment (para 5–10(a)(2)), and/or referral of employees who become ill during duty hours.

b. A medical evaluation in support of a claim controversy and for employees who are expected to be absent from work for 2 weeks or more due to a job-related illness or injury. Medical personnel will provide this evaluation by reviewing medical reports and/or performing an appropriate examination. Specialty consultation should be requested when indicated.

c. Evaluation of employee health status on return to duty after any absence due to job-related illness or injury.

d. Evaluation of employee health status on return to duty after absence due to illness not described in c above. The IMC and appropriate personnel officers will determine the duration of absences or types of illness or injury requiring such evaluation with the exception of food handlers and patient care personnel. Employees excepted will report to the OH service for evaluations after any absence due to illness.

e. Recommendations concerning work limitations.

5–14. Chronic disease and handicapped personnel

Civilian employees with chronic diseases or disabilities can be productive members of the work force. The following employment guidelines will be used:

a. Medically evaluate their work capability as a basis for proper job placement.

b. Identify employees with chronic diseases or disabilities that may affect or be affected by the work assignment. Health records will be identified and will contain clinical data regarding the condition and current treatment and the name of the personal physician.

c. Provide periodic counseling to the employee and/or supervisor when indicated.

5–15. Occupational vision

a. An occupational vision program oriented toward preservation of eyesight is an essential part of the Occupational Health Program. Guidance provided in TB MED 506 will be followed to develop and conduct the occupational vision program. An effective occupational vision program must include—

(1) Determination of which jobs or areas are eye hazardous.

(2) A job analysis to determine the visual skills required for optimal job performance.

(3) A visual assessment of workers, through use of an approved vision screening device, to determine whether they possess the required visual skills.

(4) Ocular surveillance (per OTSG Policy Letter 86–01.0) of personnel whose occupations are in the laser or microwave field and biennial vision screening for workers in all other potentially eye-hazardous job positions.


(6) Elective periodic vision screening for employees in noneye-
5–16. Hearing conservation

a. General. The hearing conservation program is designed to protect the employee from hearing loss due to occupational noise exposure. Implementation and maintenance procedures appear in TB MED 501.

b. Program functions.

(1) The installation commander will—

(a) Meet the hearing conservation program requirements according to this regulation and AR 385–10.

(b) Issue a command emphasis letter endorsing the installation’s hearing conservation program.

(c) Include hearing conservation as an item of interest in the local command inspection program.

(2) The IMA will—

(a) Ensure that a physician determines the diagnosis of noise-induced hearing loss. (See TB MED 501.)

(b) Notify the civilian personnel officer of an individual sustaining a permanent hearing loss which creates a hazard to the individual and others.

(c) Maintain audiometric testing and noise exposure records. (See AR 40–66, AR 25–400–2, and TB MED 501.)

(d) Provide audiometric test records and exposure information on request. (See TB MED 501.)

(e) Report significant threshold shift. (See chap 3 and TB MED 501.)

(f) Provide health education materials on request. (See TB MED 501.)

(3) The flight surgeon will fit the SPH-4 aviator’s helmet and inspect the helmet condition annually per AR 95–3.

(4) The safety officer (per AR 385–10) will—

a. Conduct inspections.

b. Include noise hazard abatement projects in the hazard abatement plan.

(5) The civilian personnel officer will—

(a) Ensure that OH is included on the inprocessing and outprocessing checklists for new, transferring, or terminating personnel. (This alerts the IMA of the audiometric evaluations required for these personnel.)

(b) Include (per AR 385–10) in the job description, where applicable, the requirement to wear PPE (hearing protectors).

(c) Ensure (per AR 385–10) that the following responsibilities are included in a civilian supervisor’s performance standards, where applicable—

1. Enforce the use of PPE.

2. Ensure that employees report for mandatory medical examinations.

(d) Notify supervisors when termination audiometric evaluations are required for individuals under their supervision. Include termination audiograms on the outprocessing checklists for noise-exposed personnel.

(e) Inform the IMA and safety officer of all workers’ compensation claims for hearing loss.

(f) The director of engineering and housing will—

(a) Erect and maintain danger signs per AR 385–30 and AR 420–70.

(b) Implement, whenever feasible, acoustical engineering control measures when exposures to steady noise exceed the time-weighted criteria.

(g) The hearing conservation officer will manage and coordinate all aspects of the hearing conservation program outlined in this regulation. These responsibilities include—

(a) Drafting and staffing an installation standing operating procedure (SOP) detailing the hearing conservation program.

(b) Ensuring that medically trained personnel fit individuals with preformed earplugs, and then examine individuals at least annually to ensure proper earplug condition and fit.

(c) Requisitioning and maintaining a supply of preformed earplugs.

(d) Providing a pair of preformed earplugs and carrying case to all noise-exposed personnel.

(e) Ensuring that monitoring audiometry is performed per TB MED 501 and USAEHA Technical Guide (TG) No. 167.

(f) Providing health education annually.

(g) Conducting unannounced inspections of noise-hazardous areas.

(h) Evaluating program participation, quality assurance, and program effectiveness.

(8) The industrial hygiene program manager, per TB MED 503, will—

(a) Use approved and calibrated equipment, and survey all suspected noise-hazardous areas and equipment at least once and within 30 days of any change in operations.

(b) Establish a time-weighted average for all civilians working in noise-hazardous areas and soldiers working in noise-hazardous industrial type operations.

(c) Maintain a current inventory of all noise-hazardous areas using DD Form 2214 (Noise Survey) until HHIM can accommodate noise information.

(d) Provide the names of noise-exposed personnel and the magnitude of their noise exposure to the—

1. Hearing conservation officer.

2. Unit commander or supervisor of the individual.

(e) Establish risk assessment codes (per AR 385–10) and forward the noise survey results, which indicate a violation, to the designated safety and occupational health official for inclusion in the violation inventory log.

(f) Establish appropriate contours and advise unit commanders or supervisors how to properly post these contours.
Unit commanders or supervisors of noise-hazardous areas will —
(a) Appoint a unit hearing conservation manager and ensure that this individual inspects helmets and/or noise muffs and requisitions hearing protectors to ensure an adequate supply.
(b) Prepare a unit SOP detailing the hearing conservation program.
(c) Purchase new equipment that generates the lowest noise levels feasible.
(d) Notify the IMA of any suspected hazardous-noise levels or changes in hazardous-noise levels in their work areas.
(e) Endorse a command emphasis letter explaining the importance of hearing conservation.
(f) Provide appropriate hearing protectors free of charge to their noise exposed personnel per AR 385–10.
(g) Ensure that noise-exposed personnel under their supervision —
1. Are provided appropriate audiometric evaluations.
2. Attend annual health education briefings.
3. Follow recommendations from audiometric examinations, medical evaluations, and noise surveys.
4. Wear hearing protectors.
5. Report for scheduled medical examinations.
6. Are notified of their exposure measurements.
7. Are allowed to choose from the appropriate approved hearing protectors.
(h) Ensure that all soldiers and noise-exposed civilians under their supervision retain a pair of preformed earplugs as an item of individual equipment.
(i) Require noise-exposed soldiers (per AR 670–1) to wear earplugs and carry the earplug carrying case as part of the battle dress uniform when appropriate.
(j) Provide copies of regulations, technical bulletins, and other hearing conservation documents to employees, or their representatives, on request.
(k) Ensure that noise-hazardous areas and equipment are marked with proper danger signs and decals.
(m) Monitor the use of engineering controls.
(n) Refer any personnel under their supervision to the MTF for any hearing problems or complaints associated with the wearing of hearing protectors.
(o) Initiate disciplinary action when appropriate.
(10) Noise-exposed personnel will —
(a) Correctly wear approved and properly fitted hearing protectors when exposed to hazardous-noise levels.
(b) Report for all scheduled medical examinations and health education briefings concerning hearing conservation.
(c) Report any hearing problems or difficulties associated with hearing protectors to their supervisor.
(d) Maintain hearing protectors in a sanitary and serviceable condition.
(e) Wear noise dosimeters to evaluate noise exposure, when requested.
(f) Keep hearing protection in their possession.

5–18. Occupational health education
a. The Occupational Health Education Program is an integral part of the Occupational Safety and Health Program.
b. The objectives are to —
(1) Ensure that employees (civilian and military) are aware of the actual and potential hazards of their workplace.
(2) Identify, evaluate, and modify those work practices that can be changed through OHE.

5–19. Chemical and/or nuclear surety
OH support will be provided to all workers involved in chemical and/or nuclear surety operations. The exact services will depend on the onsite exposures, but will include all medical aspects of the chemical and nuclear surety programs as described in AR 50–5, AR 50–6, and DA Pam 40–8.

5–20. Reproductive hazards
a. The reproductive hazards program assures that —
(1) Male and female employees are informed about potential work area reproductive hazards.
(2) The pregnant employee (military and civilian) and her fetus are not endangered by the employee's work assignment.
b. The program will include —
(1) Identifying work areas or occupations that present potential health reproductive hazards.
(2) Counseling all employees during preplacement or periodic job-related examination about the nature of any potential hazards to reproduction.
(3) Informing females about availability of job accommodation or transfer in the event of pregnancy (FPM chap 630 and AR 40–501).
(4) Instituting policy or procedure to ensure prompt notification to the OH clinic by pregnant employees as soon as the pregnancy is known.
(5) Assessing the employee's job assignment and work environment when pregnancy is known. When justified, specific job limitations should be recommended after consultation with the person's physician. Limitations due to pregnancy will be treated like any other medically certified temporary disability (FPM chap 630, AR 40–501, and AR 635–100).
(6) Providing periodic follow-up and counsel as indicated, including pregnancy outcome evaluation.

5–21. Records and forms
a. Obtain a health history from each permanent civilian employee upon employment and initiate a medical record. Records will be maintained by the appropriate MTF and kept confidential according to AR 40–66. They will be disposed of under AR 25–400–2. Entries into medical records will meet the requirements of AR 40–66. Medical records of civilian employees who are also military medical beneficiaries will be cross-coded to identify this dual status.
b. Results of atmospheric sampling affecting the employee conducted under the Occupational Health Program will be included in the military or civilian medical records and retained per AR 25–400–2. Documentation of sampling, even for negligible results, is important in assessing the present and past exposure history and in meeting legal obligations. (Atmospheric sampling results and recommendations will be posted in the work area to notify the employee and the supervisor.)
c. The following forms are available for use by the OH service.
(1) Outpatient treatment record forms authorized by AR 40–66.
(2) DD Form 689 (Individual Sick Slip).
(3) DD Form 1141 (Record of Occupational Exposure to Ionizing Radiation).
(4) DD Form 2215 (Reference Audiogram).
(5) DD Form 2216 (Hearing Conservation Data).
(6) Office of Workers' Compensation Program Forms under FPM chapter 810.
(7) SF 78 (United States Civil Service Commission Certificate of Medical Examination).
(8) SF 93 (Report of Medical History).
d. Nonmedical forms that may be filed in the employee medical record to provide supplementary medical data include—

(1) OF 345 (Physical Fitness Inquiry for Motor Vehicle Operators).

(2) SF 177 (Statement of Physical Ability for Light Duty Work).

e. Overprints of standard forms may be used when approved under AR 40–66 to meet specific needs; such as, a hazard specific health history check list overprinted on SF 600 (Health Record—Chronological Record of Medical Care).

Section IV
Industrial Hygiene

5–22. Reports

a. DA Form 3076 (RCS MED–20). RCS MED–20 will be submitted biannually by each MTF according to paragraph 3–12.

b. DA Form 3075. DA Form 3075 will be used according to paragraph 3–12 to assist in compiling data for RCS MED–20.

c. Log of Federal Occupational Injuries and Illnesses (unnumbered OSHA form). (See AR 385–40, 29 CFR 1900.67, and OSHA 2014 for use.) Coordination with the safety officer is required to assure complete collection and appropriate review and use of report data.

Section V
Personal Protective Equipment

5–25. General

The use of PPE is an integral part of the Occupational Safety and Health Program for all soldiers and civilian employees and requires input from both medical and safety personnel who are qualified in determining when, where, and what type of equipment will be used. Individuals who deliberately or carelessly violate regulations regarding the wearing of personal protective equipment and clothing will be subject to disciplinary action under AR 690–700, appendix A, and the Uniform Code of Military Justice (UCMJ).

5–26. Functions

a. Installation, activity, and/or unit commanders will provide PPE to persons who, by the nature of their jobs, are required to wear this equipment (AR 385–10).

b. Installation and/or activity safety personnel, with assistance from MEDDAC or MEDCEN industrial hygiene personnel, will—

(1) Designate areas requiring the use of PPE (such as eye-hazardous areas or areas requiring the use of a hard hat).

(2) Ensure that all PPE is stored and maintained properly.

(3) Ensure that all PPE is used as required.

c. MEDDAC and/or MEDCEN industrial hygiene personnel will—

(1) Conduct the OH hazard evaluation (para 5–24) to identify areas where potential exposures may require the use of PPE such as protective eyewear, respirators, or hearing protectors.

(2) Evaluate the adequacy of the following:

(a) Safety glasses and other eye-protective equipment.

(b) Earplugs and other hearing protective equipment according to paragraph 5–13 and TB MED 501.

(c) Provide technical assistance for the installation respiratory protection program per AR 11–34 and TB MED 502. Medical fitness requirements for respirator use are addressed in TB MED 509.

(d) Provide technical assistance and guidance within the installation hazard communication program.

(e) USAEHA will provide assistance to MEDDACs and/or MEDCENs and installation commanders in the selection of approved PPE upon request.

Section VI
Asbestos Monitoring

5–27. Medical surveillance

Medical questionnaires must be administered to all employees who—

a. Are exposed to asbestos, tremolite, anthophyllite, actinolite, or a combination of these minerals above the action level.

b. Will therefore be included in the medical surveillance program according to TB MED 513.

5–28. Forms

a. DD Form 2493–1 (Asbestos Exposure, Part I—Initial Medical Questionnaire) must be obtained for all new employees who are provided preplacement medical examinations according to TB MED 513. This initial questionnaire must also be obtained on all employees currently enrolled in the asbestos medical surveillance program if they have not previously completed the form.

b. DD Form 2493–2 (Asbestos Exposure, Part II—Periodic Medical Questionnaire) must be obtained for all employees who are provided annual medical examinations according to TB MED 513.

c. These forms will be filed in the civilian employee’s medical record and the military health record.

Chapter 6
Community and Family Health

6–1. General

Community and family health programs are intended to improve the level of health and increase the potential for self-sufficiency for servicemembers, their families, and other members of the military community. The cornerstone of the programs is a comprehensive community assessment that includes evaluation of health needs, based on actual or potential health problems, identification of and coordination with resource agencies, and prioritization of specific community and family health programs. A program document developed by the community health nursing section of PVNTMED service will describe the community and list nursing objectives and goals designed to promote, protect, and restore health.

6–2. Goals

The goals of the community and family health programs are to—

a. Discover and assess actual and potential health problems of persons and families.

b. Assist persons and families in understanding their health problems and how to cope with them.

c. Assist persons and families in changing their behavior or their environment to promote health and safety.

d. Provide or secure health care and other services that persons or families may need, but cannot provide for themselves.

e. Provide or secure support for persons and families in times of stress as an interim measure while they learn to resolve or accept their situation.

f. Coordinate with and use community resources for individual families to conserve and achieve maximum use of Army CHN resources.

g. Provide continuity in health care by planning and supporting the transition from hospital to home care.

6–3. Functions

a. MEDDAC and/or MEDCEN commanders, MEDDAC and/or MEDCEN commanders have overall responsibility for the health care of persons or families in the military community.

b. Chief, PVNTMED. The chief, PVNTMED, through the CHN, assists in the overall responsibility as the principal health adviser
with extensive access in the residential community and as the interface agent between the MTF and the community.

c. CHN. The specific functions of the CHN include but are not limited to—

(1) Assessing the total health needs, morbidity trends, and resources in the community.

(2) Planning, developing, organizing, implementing, and evaluating programs to meet the identified needs in the community.

(3) Providing health promotion, health education, and disease prevention programs for servicemembers and other members of the military community.

(4) Conducting a home visiting program.

(5) Supporting patient advocacy in the community.

(6) Evaluating, training, and supervising other healthcare personnel in community health nursing.

(7) Writing prescriptions for medicines when authorized by the MEDDAC and/or MEDCEN credentials committee per AR 40–2 and AR 40–400.

(8) Providing consultation to the OH nurse as outlined in paragraph 5–4h.

(9) Acting as health adviser to the CDS staff as directed by the MEDCEN or MEDDAC commander.

6–4. Program elements


(1) Maternal health is concerned with the physical and emotional health of pregnant and post partum women and the parenting role of women in general. The CHN will assure that programs are provided for the military community that will enhance the understanding of all phases of the birth process and will foster awareness of the stress associated with the role of parenting in the military community.

(2) The effects of the work environment on the pregnant employee and her fetus, and the effect of the pregnant employee on the worker and her fetus, and the effect of the pregnant employee on the military dependents.

b. Child health.

(1) Program availability. The CHN will assure that programs are available to provide parent education, well-child supervision, and support services for high-risk families. Programs will be fully coordinated with those of other military health and welfare organizations and civilian organizations.

(2) CDS support. PVNTMED personnel will provide health and environmental support to CDS staff and facilities per AR 608–10. Specifically—

(a) The CHN will provide health consultation to CDS staff regarding communicable disease control and disease prevention, assist CDS staff with “special needs” infants and children, and provide education and training in health related areas.

(b) Environmental health section staff will provide environmental consultation and inspection to center-based programs; specific services include, for example, consultation, inspection, and training to center-based programs, depending on local policy.

(3) Army family advocacy program (AFAP) support. The CHN will support AFAP per AR 608–18. Specifically, the CHN will—

(a) Provide services directed toward prevention of spouse and child abuse; for example, health education to individuals, families, and groups on subjects such as prenatal, parenting, and child development concerns.

(b) Assist with identifying high-risk families and provide direct services to selected families.

(c) Act as nursing consultant to the MTF staff to identify suspected abuse and neglect cases.

(d) Refer cases to a family advocacy case management team (FACMT) when the CHN suspects that spouse abuse or child abuse and neglect may exist.

(e) Provide nursing input into the assessment, intervention, and evaluation process of individual cases discussed during FACMT meetings.

(f) Receive referrals from FACMT for family health counseling and provide this service in the clinic, CHN office, or family home.

(g) Conduct a nursing assessment of the family in the home when indicated.

(4) School health. The Army CHN will provide health program supervision at on-post dependent schools that are not otherwise provided with public health nursing services. Close liaison should be maintained with nursing service at all schools attended by military dependents.

c. Community health. Programs will be established to identify persons or groups having special health needs and to provide and/or coordinate required health services.

d. Chronic disease control. The objectives are to—

(1) Identify risk factors that are associated with specific chronic diseases.

(2) Evaluate and control environmental factors that are associated with the development of chronic disease (such as occupational hazards and safety hazards).

(3) Promote knowledge regarding risk factors associated with the development of chronic disease.

(4) Facilitate, through education, the change of client behavior to reduce risk factors associated with the development of chronic disease.

(5) Identify persons at risk of developing chronic disease and initiate preventive and early treatment programs.

(6) Participate in a coordinated plan of rehabilitation for personnel with chronic disease.

e. Family safety. Family safety will be addressed by the CHN.

(1) As a routine part of home evaluation, the CHN will be alert to the common hazards in the home and instruct the family on safety.

(2) All home accidents resulting in medical care should be reported to the CHN.

6–5. Referrals

a. Referrals will be prepared under AR 40–407.

b. Telephonic referrals are acceptable for emergency situations or to alert the CHN of the written referral.

c. Referrals to the health nurse should be made regarding, but not necessarily limited to—

(1) All infants with a diagnosis of failure to thrive.

(2) Cases in which multifaceted health and social problems require home evaluation in planning and providing effective treatment.

(3) Cases requiring continuity of care from hospital to home.

(4) Persons or groups requiring health counseling regarding control of communicable or chronic diseases.

(5) Patients that require other health nursing services.

Chapter 7
Health Information and Education

7–1. General

Health information embodies the dissemination of information to the general public to raise awareness of good health practices, nutrition, and physical fitness; to raise the public’s index of suspicion regarding specific disease indicators; to inform the public of risk factors that will predispose them to the development of disease; and to inform the public of available health services. Health education is a more specific and individual effort to influence patients to become active participants in their treatment or health maintenance by modifying their behavior or lifestyle.

7–2. Goals

The goals of health information and education are to—

a. Inform the military community through the mass media about health, illness, and disability with general instructions on health protection, including efficient use of healthcare delivery systems.

b. Inform and motivate health practitioners to use all available health education resources for their patients. It must be understood
that health education programs are not intended to replace the patient-practitioner relationship and individual counseling.

c. Furnish a comprehensive service within the MEDCEN or MEDDAC to educate patients to assume maximum responsibility for their treatment and health maintenance applicable to their condition. Appropriate behavioral modification will improve management and reduce dependence on the MTF.

7–3. Functions
   a. TSG will provide consultation and assistance to commanders in planning and implementing programs of health information and health education.
   b. MEDCEN and MEDDAC commanders will encourage and promote health information and education programs by programming resources for the development of such programs.
   c. Chief, DPCCM, in coordination with chief, PVNTMED service, will plan, develop, implement, and evaluate programs of health information for the community and health education for the patient. The extent and the effectiveness of the program will depend on the resources obtained and resourcefulness of the personnel.

Chapter 8
Medical Safety
8–1. General
This chapter contains—
   a. The practices and procedures that govern the accomplishment of a safety program in Army MTFs and that supplement Army safety regulations in the AR 385-series.
   b. Guidance for medical commanders, MTF safety personnel, supervisors, PVNTMED personnel, OH personnel, and hospital engineering and/or maintenance personnel to provide a safe and healthful environment for the staff, patients, and visitors in an MTF.

8–2. Army Medical Department unit safety
   a. Safety program management functions and responsibilities for MEDDACs and MEDCENs are the same as for all units as prescribed in AR 385–10.
   b. Accident reporting and recordkeeping procedures and responsibilities are defined in AR 385–40.

8–3. Hospital safety
   a. The hospital commander has overall responsibility for safety in the hospital. However, all personnel who are employed in an MTF must be involved in an active safety program.
   b. An SOH advisory council committee will be organized with representation from the administration, PVNTMED service, medical staff, nursing staff, engineering and maintenance, housekeeping, and nutritional care. The committee will meet regularly and keep written minutes of its meetings. The findings of the committee and appropriate recommended corrective actions will be reported in the committee minutes and will be signed and approved by the commander.
   c. The hospital will have written safety policies to include procedures for safety of patients and accident reporting procedures.
   d. A safety officer or safety manager will have authority defined in writing to act upon hazardous conditions within the hospital.
   e. A safety orientation program will be provided for all new employees. Ongoing safety education will be provided by the supervisor for all employees and will be documented. Educational programs will be developed for specific areas and activities within the hospital. All employees will be instructed by their supervisors as to the hazards inherent in their jobs and the safety rules pertaining to their specific duties. Education related to job hazards will be coordinated with OH personnel, the safety manager or officer, and the infection control nurse, when appropriate.
   f. The hospital will have a written policy that prohibits smoking in the hospital or permits smoking only in designated areas. As a minimum, smoking will be controlled to protect the rights of nonsmokers from secondhand smoke. Smoking will be specifically prohibited in patient care areas, laboratories (including dental labs), darkrooms, supply rooms, pharmacies, dining facilities, snack bars, hospital exchanges, storage rooms, medical supply warehouses, material distribution supply areas, mechanical rooms, stairways, locker rooms, and where flammable and combustible liquids, and flammable gases and oxygen are used or stored.

8–4. Medical systems safety and health
   a. Hazards.
      (1) Significant safety and health exposures that are unique to treatment facilities can affect patients, visitors, and staff with potentially lethal consequences.
      (2) PVNTMED service personnel will develop a close working relationship with the MEDDAC safety officer or manager and engineering and maintenance personnel to coordinate the identification and elimination of hazards. Refer to USAEHA TG No. 152 for additional guidance.
   b. Fixed facilities.
      (1) Electrical safety requirements are defined in the Accreditation Manual for Hospitals (AMH), Joint Commission on Accreditation of Healthcare Organizations (JCAHO); the National Fire Prevention Association (NFPA) Standards 70 and 99; and 29 CFR 1910.
      (2) Fire safety requirements are defined in the AMH, JCAHO; NFPA Standards; and 29 CFR 1910.
      (3) General safety requirements are defined in the AMH, JCAHO; 29 CFR 1910; and the AR 385-series.
      (4) Environmental health requirements are defined in TB MED 2, the AMH, JCAHO; and chapters 4 (sec III), 11, 12, and 14 of this regulation.
   c. Field operations. Until definitive military-unique safety criteria for field operations and mobile facilities are issued, every attempt should be made to comply with the intent of codes and standards for fixed facilities (b above).

Chapter 9
Radiation Protection
9–1. General
   a. The Army radiation protection program is directed towards safeguarding personnel from unnecessary exposure to ionizing and nonionizing radiation. The program is based on—
      (1) Control of radiation sources.
      (2) Control of personnel.
      (3) Monitoring and education of personnel.
      (4) Measurements of radiation emissions.
      (5) Medical examinations to detect evidence of changes due to radiation.
   b. Protection depends on exercise of command authority, individual compliance, and an organization specifically devoted to radiation protection.

9–2. Purpose
This chapter prescribes the methods for the control of potential health hazards resulting from the procurement, possession, storage, transportation, use, and disposal of radioactive materials or equipment capable of producing potentially hazardous ionizing or nonionizing radiation. This chapter is not intended to conflict with or supersede established procedures for radiological defense.

9–3. Goal
The two radiation protection primary goals are to minimize the—
   a. Ionizing radiation exposure (individual and collective) to as low as reasonably achievable (AR 40–14/DLAR 1000.28).
b. Release of radioactive effluents into the environment (AR 200–1).

9–4. Organization

a. Radiation control committee. AR 40–14/DLAR 1000.28, AR 385–11, and TB MED 525 prescribe the requirements for the composition and responsibilities of the radiation control committee. The radiation control committee should also advise the commander on nonionizing radiation hazards and methods to control these hazards.

b. Radiation protection officer.

(1) The commander will designate (in writing) an RPO and an alternate RPO whose primary duties are to manage the radiation protection program. The RPO will be provided training, equipment, and a support staff commensurate with the extent of his or her responsibilities. The RPO will be responsible for managing the ionizing radiation protection requirements specified in AR 40–14/DLAR 1000.28, AR 385–11, and TB MED 525 as well as the Nonionizing Radiation Protection Program (AR 40–46 and USAEHA TG No. 153). Complete program files should be maintained by the RPO to include the current records of inventory, SOPs, and records of related safety instruction.

(2) Medical activities with nuclear medicine services require a full-time RPO qualified under TB MED 525.

9–5. Medical surveillance

a. Personnel potentially exposed to ionizing radiation in their occupational environment will receive medical examinations as required by AR 40–14/DLAR 1000.28.

b. Personnel potentially exposed to nonionizing radiation will receive medical examinations as required by AR 40–46, TB MED 523, TB MED 524, and USAEHA TG No. 153.

9–6. Personnel dosimetry

a. Ionizing radiation.

(1) The DA policies regarding ionizing radiation exposure standards, the use of personnel dosimeters, and the recording of occupational exposure to ionizing radiation are specified in AR 40–14/DLAR 1000.28.

(2) Bioassay procedures will be performed when radioactive materials are used in such a manner that they could be inhaled, ingested, or absorbed into the body (AR 40–14/DLAR 1000.28).

(a) The necessity, frequency, and methodology for performing bioassay procedures will depend on the radionuclide(s), their chemical and physical form, and the amount of material potentially available for entry into the human body (AR 40–14/DLAR 1000.28).

(b) Unless prescribed elsewhere, the type of analysis and the frequency of the bioassay procedure will be determined by the IMA in consultation with the RPO.

(c) Bioassay services are available from the Commander, USAEHA, ATTN:HSHB–ML–R, Aberdeen Proving Ground, MD 21010–5422. Specimens should be sent directly to USAEHA. Do not send the specimens through command channels. Each specimen should be accompanied by a properly completed laboratory form (SF 557 (Miscellaneous)). The following information, in addition to the data normally entered on laboratory forms, is required in the remarks block to enable USAEHA to evaluate the internal exposure from radioactive material:

1. Possible exposure date and time (if known), in case of acute exposure.
2. Duration of exposure, in case of chronic exposure.
3. Chemical and physical form of radionuclide and suspected activity involved.
4. Suspected route of entry.
5. Any additional information concerning the situation that prompted the request (if necessary, on a separate sheet or consultation sheet (SF 513, (Medical Record—Consultation Sheet)).

(3) All actual or alleged overexposures to ionizing radiation will be investigated and reported under AR 40–14/DLAR 1000.28 and AR 385–40.

(4) The RPO will calculate the collective exposure to ionizing radiation of all persons for which a DD Form 1141 is maintained. This calculation will include the most recent 3 months of reported exposures.

(5) The RPO will report quarterly the following information to the radiation control committee:

(a) Collective exposure: Person-roentgen equivalent man/person-rem in a quarter. (Person-rem is calculated by adding all exposures during a quarter.)

(b) Average exposure: rem/quarter. (Rem/quarter is obtained by dividing person-rem by the total number of persons monitored.)

(c) Highest exposure: rem.

b. Nonionizing radiation. No suitable personnel dosimeters are available.

9–7. Protective clothing and equipment

a. Ionizing radiation.

(1) Concentrations of airborne radioactive materials located in the breathing zone of radiation workers will not exceed concentrations as specified in 10 CFR 20.103 without adequate respiratory protection (TB MED 502).

(2) Protective clothing and respiratory protective equipment may be required to minimize the exposure of the worker. When required, such equipment and clothing will be identified for control purposes. (See AR 385–10 and AR 385–32.)

(3) Adequate respiratory protection programs will be established to assure that equipment appropriate to the potential hazard is selected, used, and maintained (TB MED 502). The use of respiratory protection is not a suitable substitute for proper ventilation, containment, and process controls, and will not be used in place of other feasible engineering controls.

(4) A respirator that is not used routinely, but maintained ready for emergency use, will be inspected after each use and at least monthly to assure that it is in satisfactory operating condition (TB MED 502). A record will be maintained of inspection dates and findings (AR 25–400–2).

(5) When laboratory hoods are used to maintain minimum levels of airborne radioactive material in work or storage areas, the airflow in the hood will have an average velocity of at least 100 linear feet per minute (fpm) (30 meters/minute (m/min)) plus-or-minus 10 percent through the fully open face. Glove box hoods must have an inward average velocity of 50 fpm through doors/ports or 0.25 inch static pressure on a closed system. Hoods should be provided with a dual speed fan to permit operation at a higher velocity while the hood is in use and at a lower velocity when it is closed. Bypass openings will be provided to maintain proper hood and room pressure balance. The variations in air velocity through the open face will not exceed plus-or-minus 20 percent. Each hood will have an independent exhaust system with the fan installed outside the building or at the point where the exhaust leaves the building to ensure that the duct work inside the building is under negative pressure. The point of discharge should be at least 10 feet (3.1 m) above the roof and 100 feet (31 m) from any air intake to minimize radioactive effluents being carried back into the same or adjacent buildings. The fan should discharge into a vertical stack with no directional baffles or projections (TM 5–810–1).

(6) Laboratory hoods should be evaluated and flow measurements made at least semiannually. Documentation of such measurements will be maintained under AR 25–400–2.

b. Nonionizing radiation. PPE is only used when other control measures do not provide adequate protection. TB MED 524 provides guidance for the proper use and marking of laser eye protectors.

9–8. Radiation detection and measuring equipment

a. Ionizing radiation.

(1) All radiation protection survey instrumentation for health and safety monitoring of radiation sources will be calibrated at the frequency specified in TB 750–25.

(2) All pocket dosimeters used for health and safety purposes will be calibrated, inspected, and certified.

(3) Radiation protection survey instrumentation will have a DA
The operational status of all radiation protection survey instrumentation will be verified before removal of shielding from a radiation source or entering a known or suspected radiation field. All radiation measuring equipment used with nuclear medicine procedures will be calibrated as specified in TB MED 525. All dosimetry systems used in the calibration of therapeutic x-ray and gamma-beam equipment will be calibrated as specified in TB MED 521.

b. Nonionizing radiation. Other than the use of recommended instrumentation for the measurement of microwave oven leakage, the instrumentation required to adequately assess the hazards of high intensity ultraviolet, infrared, visible, ultrasound, radio frequency, microwave, and laser radiation is highly specialized and expensive. Therefore, it should only be purchased when personnel capable of making the assessment are assigned.

9–9. Control of radiation sources

a. Control of both ionizing and nonionizing radiation hazards. (1) Commanders of installations or activities responsible for the design or development of equipment capable of producing radiation or the incorporation of equipment or radioactive materials into Army systems will ensure that such equipment or devices have been evaluated for potential health hazards by USAEHA. This evaluation will take place during the research, development, test, and evaluation phase of the equipment and before acceptance or adoption (AR 70–1 and AR 385–10). A reevaluation of the equipment must be made if substantial modifications are made between the initial USAEHA evaluation and final acceptance or adoption.

(b) Procedures to be followed when an accident or incident occurs or in other emergency situations.

1. Safe working techniques and procedures.
2. Proper use of protective equipment and devices.
3. Procedures to be followed when an accident or incident occurs or in other emergency situations.
4. Daily preoperational, operational, and postoperational checks or surveys to ensure proper radiation safety.
5. Procedures for maintaining an operational log for each piece of equipment that will identify when interlocks and other control or warning devices are bypassed or overridden.
6. Records of these instructions will be maintained by the RPO. They will include a brief outline of the instructions and a list of persons who received these instructions (AR 25–400–2).
7. All controlled areas will be properly marked, have proper warning signs, and, where required, have proper warning signals and safety switches (AR 385–30, TB MED 521, and TB MED 525).
8. Individuals are designated (in writing) to receive notice in the event of emergencies such as major spills or accidental release of radioactive material, bodily injury, fire, and major malfunction of equipment that may produce or generate potentially hazardous radiation fields. A list of those persons and phone numbers will be posted in each controlled (restricted) area.
9. A comprehensive inventory of radioactive material and equipment capable of producing radiation will be maintained (AR 40–46, AR 385–11, and TB MED 525). A copy of this inventory will be forwarded to the RPO.

b. Control of ionizing radiation hazards. (1) Commanders of installations or activities will not possess, use, or transfer radioactive materials or use equipment capable of producing ionizing radiation in such a manner that could cause any person to receive a dose equivalent in excess of the radiation exposure standards specified in AR 40–14/DLAR 1000.28. Personnel responsible for the Radiation Protection Program will be continually vigilant concerning means to reduce exposure of personnel to ionizing radiation.

(2) The prevention of radioactive contamination is especially important for persons working with unsealed radioactive materials. Every user will maintain constant vigilance to minimize or prevent contamination and to contain its spread (AR 385–11 and TB MED 525).

(3) The RPO will perform surveys of all laboratories and work areas where radioactive materials are used or stored (AR 385–11, AR 700–64/DLAM 4145.8/NAVSUPINST 4440.34/MCO 10330.2/AFR 67–12, and TB MED 525). Radiation surveys should be performed routinely to indicate any changes in radiation levels in the work area. The surveys will evaluate the effectiveness of controls and procedures, ventilation, respiratory protective equipment, fixed and transferable surface contamination, airborne radioactive materials, and general exposure levels in the work area and environment. The frequency of any radiation survey will depend on such factors as the type of operation, the type and level of the radiation, the rate at which changes could unknowingly develop, the potential hazard, and the degree of personnel involvement. Since there may be possibilities of radiation or radioactive contamination occurring in generally unexpected locations, it is desirable to occasionally monitor so-called “clean/cold” or uncontrolled (unrestricted) areas.

(4) Smoking, eating, drinking, or applying cosmetics will not be permitted in work areas where unsealed radioactive materials are used or stored. Food or drink will not be stored in an area where radioactive materials are stored.

(5) To reduce the possibility of fire or other major disasters, buildings where radioactive materials are used and stored should be constructed of fire retardant materials. Fire prevention and military police personnel will be informed of any buildings or areas where potential radiation hazards may exist. Specific conditions under which it is safe to handle emergencies will be explained carefully to firefighters, security guards, and military police, and included in the appropriate SOP. These conditions will be respected unless they are modified by the responsible RPO or safety manager.

(6) All ionizing radiation accidents or incidents will be investigated and reported according to the requirements in AR 40–14/DLAR 1000.28 and AR 385–40.

(7) The planning, procurement, installation, calibration, preventive maintenance, evaluation, and use of diagnostic and therapeutic x-ray and gamma-beam equipment will be according to TB MED 521.

(8) Radiation therapy equipment used for human treatment will be calibrated by a qualified expert as required by TB MED 521. Evidence of calibration will be retained so it will be available to a surveyor or inspector (AR 25–400–2).

(9) As required by TB MED 521, a radiation protection survey by a qualified expert will be performed on all new or modified diagnostic or therapeutic facilities before clinical use.

(10) Industrial radiographic facilities will be classified and governed by procedures or conditions of the facility’s Nuclear Regulatory Commission (NRC) license, DA radioactive material authorization or permit, or National Bureau of Standards Handbooks (NBSHs) 107, 111, or 114, as appropriate.

(11) A radiation protection survey by a qualified expert will be
performed on all new or modified industrial radiographic facilities before placing the equipment in routine operation.

(12) Radiation protection surveys will be performed periodically by the RPO to determine the exposure or exposure rate in the environment during operation of the equipment. These surveys will be conducted in areas as deemed necessary to evaluate and control the potential radiation hazard (AR 385–11 and USAEHA TG No. 153).

(13) All radioactive material, other than nuclear weapons, will be transported (shipped and received) according to the requirements of AR 385–11, Technical Manual (TM) 55–315, and TM 55–4470–400–12–1, as appropriate. The reporting of packaging and handling deficiencies will be under AR 700–68/DLAR 4145.25/NAVSUPINST 4000.34/AFR 67–8/MCO P4400.105, and discrepancies in shipment will be reported under AR 55–38/NAVSUPINST 4610.33/AFR 75–18/MCO 4610.19/DLAR 4500.15.

(14) The disposal of unwanted radioactive material will be per AR 385–11.

c. Control of nonionizing radiation hazards. Commanders of installations or activities responsible for the operation or testing of nonionizing generating equipment will take the necessary measures to ensure that—

(1) The potentially hazardous system has been evaluated by USAEHA before operation, and hazards criteria for the system are available and being observed.

(2) Personnel working in the vicinity of such equipment are informed of potential personal health hazards.

(3) SOPs are published and enforced to deal with operational limitations placed on the equipment and control of the radiation field to minimize personnel exposure.

(4) Periodic operational checks are conducted on all radiation safety devices such as alarms, lights, and interlocks installed on or near radiating sources. Such tests should be conducted prior to system operation. Defective devices should be repaired or replaced before continuing operation. A log should be maintained of all such cases.

(5) Safety procedures prescribed in TB MED 523 or TB MED 524, as applicable, are being complied with.

(6) When interlocks and other control or warning devices are bypassed or overridden, operational logs must indicate the purpose and duration.

(7) All nonionizing radiation areas are properly marked, have appropriate warning signs and, where required, have proper warning signals and safety switches.

(8) All alleged overexposures or accidents involving nonionizing radiation will be reported under the requirements in AR 40–400 and AR 385–40.

d. Monitoring personnel. Monitoring personnel will ensure that personnel potentially exposed in their occupations to ionizing radiation or radioactive materials are appropriately monitored. (See para 9–5.)

9–10. Licenses and authorizations

a. TB MED 525 gives policies and procedures for the control and use of radioactive materials for medical purposes. It also prescribes the requirements and procedures for obtaining an NRC license or DA radioactive material authorization for the possession and use of materials not under specific licensing control of the NRC, the materials being used for medical purposes.

b. AR 385–11 prescribes policies and procedures for the control and use of radioactive materials for nonmedical purposes. It also prescribes the requirements and procedures for obtaining an NRC license or DA radioactive material authorization or permit for the possession and use of materials not under specific licensing control of the NRC for nonmedical purposes.

c. AR 385–11, AR 700–64/DLAR 4145.8/NAVSUPINST 4000.34/AFR 67–8/MCO 4400.105, and TB MED 522 prescribe policies and procedures for the control of radioactive commodities within the DA supply system.

9–11. Radiologic facility shielding analysis

a. Design plans for the modification of existing medical radiographic facilities and design or construction specifications for new medical radiographic facilities must be reviewed by a qualified expert prior to modification or construction.

(1) The CONUS qualified expert review will be provided by either a MEDCEN having an assigned nuclear medical science officer or the USAEHA.

(2) The OCONUS qualified expert review will be provided by either the 7th Medical Command or the USAPACEHEA.

b. Plans and design specifications for industrial radiologic facilities will be reviewed and evaluated by a qualified expert before the modification or construction of a new industrial radiologic facility.

9–12. Laser and radiofrequency radiation exposure incidents: reporting procedures and registry maintenance

a. The radiological hygiene consultant to TSQ will request that USAEHA conduct an on-site investigation when—

(1) An employee’s lesion or ocular complaint may have resulted from exposure to nonionizing radiation.

(2) An exposure to radiofrequency radiation is five times or more the permissible exposure limit.

b. If an alleged laser or radiofrequency radiation exposure occurs, the affected installation—

(1) Calls USAEHA within 24 hours to forward incident information.

(a) During duty hours, contacts the Chief, Laser Microwave Division (AUTOVON 584–3932), or the Chief, Occupational and Environmental Medicine Division (AUTOVON 584–3534).

(b) During non-duty hours, contacts the duty officer, or noncommissioned officer-of-the-day (AUTOVON 584–4375).

(2) Ensures that the potentially exposed individual(s) receive(s) appropriate medical evaluation within 24 hours of the exposure.

(3) Develops and transmits an—

(a) RCS MED–16 report per AR 40–400.

(b) RCS DD–R & E (AR) 1168 (Radiological Incident Report) per AR 385–40.

c. USAEHA will conduct investigations of alleged laser or radiofrequency radiation exposures and maintain the U.S. Army Laser and Radiofrequency Radiation Incident Registry.

Chapter 10 Pest and Disease Vector Prevention and Control

10–1. General

DA pest management, as a single comprehensive program, encompasses—

a. Personnel training.

b. Pest surveillance.

c. Recommendations for and implementation of IPM practices.

d. Assessment of environmental, safety, and health consequences of IPM practices.

e. Technical support.

10–2. Objectives

The comprehensive IPM program—

a. Provides prevention and control of pests that could cause major medical or economic harm.

b. Protects personnel and the environment from the toxic effects of pesticides.

c. Assures the preparedness of field units to prevent and control vector-borne disease in time of war, military conflict, or national disaster. (See chap 14.)

10–3. Functions

a. Armed Forces Pest Management Board. Under AR 10–64/OPNAVINST 6700.2/AFR 160–29/MCO 5420.18A, the board is a joint activity of DOD that—
(1) Develops and recommends policy to the Assistant Secretary of Defense(Production and Logistics) for the DOD Pest Management Program.

(2) Serves as a scientific advisory body to DOD components on pest management.

(3) Coordinates and develops requirements for pest management related research, development, and testing within DOD.

(4) Operates the Defense Pest Management Information Analysis Center.

(5) Maintains liaison with other Federal agencies having similar programs.

b. Installation medical authority. The IMA will—

(1) Establish and evaluate health criteria and standards of pest management programs; provide advice and technical guidance to program operators on safe storage, use, and disposal of pesticides; provide training of personnel involved in pest and disease vector control operations; and review installation and field unit pest management programs and plans, and provide technical assistance and evaluation of the health, safety, effectiveness, and environmental soundness of the programs.

(2) Conduct surveillance of vectors and pests affecting the health and welfare of the Army community; conduct ecological and biological studies of pests, as required; determine pest resistance to pesticides; monitor pesticide levels in the environment; and evaluate the safe use of pesticide dispersal equipment and material.

(3) Recommend that the commander assure the proper use of insect or arthropod repellents by soldiers entering areas where the risk of vector-borne diseases and/or troublesome numbers of pest bites occur. Maximum repellent protection can be provided to the soldier by use of the new topical repellent lotion(national stock number 6840–01–284–3982) on exposed skin surfaces and use of the clothing repellent, permethrin, on the battle dress uniform.

(4) Conduct operations, as required, for the control of disease vectors and animal reservoirs through TOE PVNTMED teams and new clothing repellent has a regulatory agency label pending. The number 6840–01–284–3982) on exposed skin surfaces and use of the clothing repellent, permethrin, on the battle dress uniform.

(5) Assure, through coordination with the facility engineer, that the installation pest management program complies with those laws and regulations relevant to the health and safety aspects of pest management.


d. Quartermaster laundry and bath units. Provide repellent impregnation of uniforms and deousing operations when recommended by the IMA.

e. USAEHA. The USAEHA will—

(1) Provide consultative, field, and laboratory services for second- and third-level support of the IMA functions listed above.

(2) Function as executive agent for the DOD pesticide regulatory action system which includes—

(a) Providing pest and pesticide information to medical and other DOD activities via the DOD Pesticide Hotline.

(b) Reviewing pest management regulatory documents proposed by Federal and State agencies.

(c) Drafting DOD responses to regulatory proposals.

10–4. Pesticides

a. The use of pesticides to control medically and economically important pests will remain an integral part of the DA pest management program. However, nonchemical IPM techniques will be emphasized to minimize the unnecessary introduction of toxic substances into the environment. Where nonchemical IPM methods fail to adequately manage a pest population, chemical control may be used as a supplement. However, chemical control will only be used in conjunction with nonchemical control measures.

b. The use of preventive or scheduled periodic pesticide treatments is prohibited unless approved by the pest management consultant concerned and based upon surveillance data or past pest problems.

c. Pesticide monitoring programs will be established to assess possible adverse environmental or public health effects and to monitor the health and safety of persons occupationally exposed to pesticides. (See para 11–10.)

d. Procurement, storage, use, and disposal of pesticides will be under AR 200–1, AR 420–76, Executive Orders, and Federal laws and regulations as applicable. Oversea commands will abide by applicable U.S. Code sections and Status of Forces Agreements.

e. Only personnel trained and/or certified under AR 420–76, and/or the DOD plan for certification of applicators of restricted use pesticides, will apply or supervise the application of pesticides.

f. Units authorized to use pesticides will procure only those items and amounts authorized in SB 3–40 and CTA 50–970. Every effort will be taken to maintain the integrity and operational usefulness of those pesticide items subject to deterioration or degradation. Medical units requiring nonstandard pesticide items or standard pesticides not identified in the TOE or TDA will submit requisitions to the MACOM surgeon for approval prior to the submission of the requisitions to supply channels.

(g. Pesticides and equipment issued or distributed to military personnel and occupants of family housing for use in self-help programs (AR 420–22) will normally be restricted to standard issue items identified by the AFPMB as being available for use by untrained and uncertified personnel. Center managers will be guided by MACOM professional pest management personnel and the IMA in standard or nonstandard item selection and issue.

h. Guidelines for selecting, selling, and handling pesticides in post exchanges and commissaries are as follows:

(1) Pesticides selected for sale in post exchanges and commissaries will be registered for “General Use” by the EPA or by the State in which the facility is located. Items with labels indicating that only professional pest management personnel may utilize the product or items labeled“Restricted Use” will not be sold.

(2) All pesticides sold in post exchange and commissary facilities will be arranged separately on sales display shelves and in storage according to type (that is, herbicides, insecticides, rodenticides, fungicides, and disinfectants). Pesticides will be segregated from all food products in storage, during transportation, and while on display. Segregation means there will be sufficient space between pesticides and food items so that spillage or leakage will not contaminate food.

(3) Employees handling pesticides will be familiar with proper measures for safe handling. They will use display practices that minimize accidental breakage by customers and the handling of products by children. They will also be familiar with cleanup procedures for spills (app C).

i. Guidelines for pesticides spills are as follows:

(1) Guidance on the cleanup and disposal of small quantity pesticide spills (1 quart (qt) or less) is given in appendix C. Larger spills will be reported and decontaminated according to the installation’s oil and hazardous substances spill control and contingency plans (AR 200–1, paras 8–10 and 8–11).

(2) Immediate assistance for emergency type pesticide spills that threaten life or gross contamination of the environment can be obtained by calling the Chemical Transportation Emergency Center (CHEMTREC). The telephone number for CHEMTREC is (800) 424–9300. For spills outside CONUS or within the Military District of Washington, call (202) 483–7616.

(3) Information on decontamination of nonemergency type pesticide spills may also be obtained by dialing the CHEMTREC number given above. The operator must be told immediately that—

(a) No emergency exists.

(b) The call is a request only for decontamination information.

j. Information on applicable Federal and DA regulations, EPA, and State registered pesticides and pesticide label information may be obtained from USAEHA by calling the DOD Pesticide Hotline at (301) 671–3773 or AUTOVON 584–3773. An answering device is available to receive questions outside the normal USAEHA duty hours.
10–5. Pest control equipment and devices
a. Equipment and devices listed in Federal Supply Catalog Identification Lists(C–3000/6300 and C–6700/9500) will normally be used in pest management operations. Only standard items will be requisitioned by medical TOE units. Activities other than TOE organizations authorized to conduct pest management operations or evaluations may requisition standard or nonstandard pesticide dispersal equipment. Requests for approval to purchase nonstandard items will be submitted to the MACOM pest management professional for approval. Approval will be obtained prior to initiating procurement action.
b. Medical personnel conducting health assessments of pest management programs will ensure that equipment used is compatible with the pesticide formulation being applied and that equipment is properly calibrated. USAEHA will provide technical assistance in the calibration of ultra low volume equipment and droplet size determination.
c. Dispersal equipment will be rinsed before the equipment is used for application of another pesticide or at the conclusion of each day’s activities. Rinse water from spray equipment will not be placed into a sanitary sewer, but should be used as a diluent for subsequent spraying operations or treated as a pesticide-related waste and disposed of per current Federal, State, or host country requirements.

10–6. Training and certification
a. Commanders will ensure that only personnel trained and/or certified under AR 420–76 and/or the DOD plan for certification of pesticides/applicators for restricted use pesticides will apply or supervise the application of pesticides except for those pesticides that have been identified specifically for use by untrained, uncertified personnel by the AFPMB (DOD Directive 4150.7).
b. The Academy of Health Sciences (AHS), U.S. Army, will conduct certification and recertification training to meet the standards outlined in the DOD plan. In addition, AHS will conduct other resident and nonresident training as requested by the Office of the Chief of Engineers and other military or Federal organizations to meet specific pest management training needs. Pest management professionals requiring certification may direct their requests to the AFPMB. Recertification must be obtained every 3 years.
c. Training and/or certification for the aerial application of pesticides is provided by the Air Force 907th Tactical Airlift Group Spray Branch, Rickenbacker Air National Guard Base, Ohio, AUTOVON 950–4694/3106 or commercial (614) 492–4694/3106.

10–7. Pest surveillance
a. Installation pest management programs will involve—
   (1) Surveillance of pest populations.
   (2) Recommendations of measures to control pests.
   (3) Application of pest control measures.
   (4) Reevaluation of pest populations to determine the effectiveness of control.
b. The IMA will conduct surveillance of pests of medical importance using the guidance provided in USAEHA TG No. 102 and recommend appropriate IPM measures. IPM recommendations will be coordinated with MACOM pest management professionals.
c. Surveillance will be recorded using the guidance provided in USAEHA TG 102. The time or labor expended by the IMA in surveillance activities will be transmitted to the facilities engineer for inclusion on the monthly DD Form 1532 (Pest Management Report). Results of surveillance will be reported to the director of facilities engineering pest management personnel for their use in determining appropriate control measures. Chemical control measures will be initiated only after nonchemical control methods have been fully implemented and have failed to control the pest population. At no time will pesticides be applied in a food handling facility without current surveillance data documenting a pest infestation.
d. Records of surveillance activities and pest management recommendations will be maintained to provide documentation and permit short- and long-term assessments of the effectiveness and environmental consequences of the installation pest management programs.
e. The USAEHA will provide pest identification services as required.
f. Upon request, USAEHA will provide special onsite investigative services in the identification and assessment of pest problems.

10–8. Pest resistance
USAEHA will support CONUS installations with pest resistance surveys and/or provide laboratory and technical assistance as required. OCONUS installations will be supported by medical laboratories in overseas commands. Pest resistance support will be coordinated with installation professional pest management personnel.

10–9. Pest management in Army food handling establishments
a. General. IPM program principles and measures will be applied in food handling establishments (as defined in 38 FR 21685). At no time will pesticides be applied in a food handling establishment without current or historical surveillance data documenting the pest infestation. The effectiveness of chemical control measures will be assessed by the IMA, and the results will be conveyed to the pest control activity. Additional guidance on pest management operations in food handling establishments is provided in TB MED 530.
b. Food preparation areas. Pesticides will be applied with food service personnel to ensure the safety, effectiveness, and efficiency of the pesticide treatment. Pesticide treatments will be conducted only when the food preparation area is not in operation and must be used according to the pesticide label precautions. Automatic aerosol pesticide dispensing devices will not be used in food serving or preparation areas. Insect electrocutors or sticky fly papers may be located in nonfood areas of food handling establishments, provided that their use is in a manner that will preclude contamination of any food or food-contact surfaces and their use is not in lieu of proper sanitation. Pesticides, except disinfectants, will not be stored in food serving facilities.

10–10. Pest management in Army MTFs
a. General. The management of pest infestations in MTFs will be accomplished through timely surveillance and effective pest management procedures according to USAEHA TG 106. MEDDAC or MEDCEN commanders will investigate pest problems and determine pest management measures required in patient areas.
b. Pesticide applications in patient areas. Pesticides will not be applied while patients are in the immediate area. Patient areas (including, but not limited to, emergency rooms, examining rooms, wards, operating rooms, infant nurseries) will not receive preventive pesticide treatments but will be treated only when an actual infestation is evident and nonchemical control measures have failed.
c. Food service areas in an MTF. Pest management in food service areas is addressed in paragraph 10–9. Cockroach infestations in portable food carts are difficult to control. Routine procedures for either nonchemical or chemical control will consist of numbering carts and subjecting them to the treatment of choice on a regularly scheduled basis (USAEHA TG 106). Carts treated with nonresidual pesticides will not be used to transport food to patients until the carts have been steamcleaned or sterilized. Residual insecticides will never be applied to food carts because of the potential of contaminating food items. Technical assistance in controlling cockroaches in food carts may be obtained from USAEHA.

10–11. Pest management of military subsistence and stored materials
The surveillance and control of arthropods, rodents, birds, and other pests of subsistence and stored products will be conducted under DOD Directive 4150.7 and MIL STDs 904 and 1486. Reporting of losses due to pests will be conducted as outlined in DA Pam 40–17. Additional technical information concerning surveillance and control of stored product pests can be found in TM 5–632/NAVFAC MO–310/AFM 9–16. Upon request, professional assistance and
guidance will be provided by the IMA or installation veterinary personnel.

10–12. Aerial dispersal of pesticides
Aerial dispersal of pesticides will be conducted under AR 40–574/AFR 91–22 and AR 420–76. All aerial dispersal of pesticides must receive appropriate prior MACOM approval. Upon request, entomologists assigned to the IMA, MACOM, or USAEHA will provide assistance in the preparation and evaluation of aerial applications. Actual application will be conducted under the direct and continuing supervision of an applicator certified in the category of aerial dispersal of pesticides.

10–13. Pesticides and pesticide container disposal
Pesticides and containers that are excess because of unserviceability, registration cancellation, or other reasons will be disposed of under 40 CFR 165. Guidance is available in AFPMB Technical Information Memorandum (TIM) No.21. In no event will pesticides or pesticide containers be disposed of in such a way as to needlessly contaminate the environment or in a manner inconsistent with the label. Contact USAEHA, Waste Disposal Engineering Division, for pesticide disposal instructions (AUTOVON 584–2024).

10–14. Protective clothing and equipment
Responsibilities, policies, and procedures for providing protective clothing and equipment are specified in AR 385–10 and TB MED 302. Items will be requested and maintained under AR 710–2. Supply management data necessary to request standard pest management protective equipment may be obtained from AFPMB TIM No. 14. Current information on protective equipment, to include respirators approved for use with pesticides, may be obtained from the Industrial Hygiene Directorate, USAEHA (AUTOVON 584–3946).

10–15. Medical surveillance
All personnel known as pesticide applicators will be included in medical surveillance, health education, and respiratory protection programs as part of the Occupational Health Program.

10–16. Pest quarantine
The USPHS and the U.S. Department of Agriculture (USDA) policies and procedures in AR 40–12/SECNAVINST 6210.2/AFR 161–4 will be followed to prevent the introduction and dissemination of animal or plant pests of medical and agricultural importance. Guidance on materials and methods suitable for quarantine operations can be found in TM 5–632/NAVFAC MO–310/AFM 9–16.

10–17. Retrograde material treatment
Retrograde programs are essential to prevent the importation into the United States, its territories, trusts, and possessions of pests of medical or agricultural importance. The IMA responsible for the operation of retrograde programs will ensure the full implementation of AR 700–93.

10–18. Personal-use protective measures
a. Commanders will assure the proper use of insect or anthropod repellents by soldiers entering areas where the risk of vector-borne diseases and/or troublesome numbers of pest bites occur.

b. Insect bed nets and head nets should be considered for use in situations where their use will provide protection from bites of vector and pest species.

c. Commanders will assure the proper wearing of the battle dress uniform by soldiers to provide protection from bites of vector and pest species.

d. DA Poster 40–5 should be placed in residential, military training, hunting, and recreational areas and wild game reserves to—

(1) Warn individuals that they are entering an area infected with ticks that can transmit tick-borne diseases such as Lyme.

(2) Explain the precautions an individual should take when entering and exiting a tick infested area.

e. DA Poster 40–5 is available through normal publications channels; however, its use is controlled as follows:

(1) CONUS—Each MEDCEN commander will decide when and where in the MEDCEN’s geographical area the poster will be used.

(2) OCONUS—Each major medical command commander (for example, 7th Medical Command Commander) will decide when and where in the command’s geographical area the poster will be used.

(3) Modification of the poster—None is allowed.

(4) Local reproduction—None is allowed.

Chapter 11
Environmental Quality

11–1. General
a. The AMEDD fulfills a major role in the Army’s effort to protect the natural environment. This chapter prescribes policies and assigns responsibilities for associated PVNTMED programs.

b. The goal is to minimize adverse environmental impact and protect community health with minimum impairment of the Army’s mission or readiness.

c. Technical support may be obtained from USAEHA, 10th Medical Laboratory, and USAPACEHEA if it is beyond the local preventive medicine organizations’ capability to provide the assistance.

11–2. Functions
a. The MACOMs—

(1) Establish procedures to ensure that health and welfare factors are considered during environmental assessments.

(2) Monitor health and welfare aspects of Army operations to ensure that they meet environmental objectives.

b. HSC, in addition to its MACOM functions, will—

(1) Ensure that PVNTMED personnel coordinate with environmental personnel to establish installation level PVNTMED liaison with local health regulatory agencies and supported Army activities and units.

(2) Provide MACOMs with technical assistance that includes investigations, consultations, special studies, and routine environmental surveys.

(3) Provide, through PVNTMED, AMEDD support of the goals, standards, and procedures issued in AR 200–1.

(4) Provide AMEDD support of the Army environmental compliance achievement program operated by the Assistant Chief of Engineers.

11–3. Wastewater
a. Objective. The DA objective is to dispose of Army waterborne wastes in a manner that protects water resources from contamination and preserves the public health.

b. Functions. PVNTMED will—

(1) Monitor environmental impact by reviewing wastewater data generated by routine and special Army surveillance programs.

(2) Monitor the performance of pollution abatement facilities by conducting frequent visits during routine operations.

(3) Assist installations in preparing applications for Federal and State discharge permits, review permits for general acceptability of specific parameters, and provide guidance for integrating the Best Management Practice Plan into National Pollutant Discharge Elimination System permits.

11–4. Air quality
a. Objective. The DA objective is to reduce pollutant emissions from Army stationary and mobile sources to the lowest practicable limit to protect health and ensure compliance with appropriate Federal, State, and local regulations.

b. Functions. Installation PVNTMED personnel will—

(1) Evaluate proposed and existing stationary and/or regulated sources of air pollution to ensure that they do not present a potential for adverse health effects.
11–5. Solid waste
   a. Objective. The DA objective is to manage Army solid waste to ensure compliance with appropriate Federal, State, and DA regulations in a manner that permits maximum opportunity for resource recovery without jeopardizing natural resources or health and the environment.

   b. Functions. Installation PVNTMED personnel will—
      (1) Monitor the installation’s management of solid wastes, including the stages of segregation, storage, transportation, and disposal and/or sale.
      (2) Evaluate community complaints and provide health and welfare recommendations to the facilities engineer.
      (3) Assist installations in preparing applications for landfill permits, review permits for general acceptability, and provide guidance in site selection.

11–6. Hazardous wastes
   a. Objective. The DA objective is to manage Army hazardous waste to ensure compliance with appropriate Federal, State, and DA regulations in a manner that permits maximum recovery and protects health and the environment.

   b. Functions. Installation PVNTMED personnel will—
      (1) Provide technical assistance regarding potential health effects for identifying unknown waste and for selecting and/or evaluating storage, treatment, and disposal methods.
      (2) Assist installations in preparing permits for hazardous waste treatment, storage, and disposal facilities; in reviewing permits for health-related implications; and in providing guidance on the operation of hazardous waste facilities in conformance with regulatory requirements and in a manner protective of health and environment.
      (3) Assist and advise hazardous waste generators on means to reduce the amount of waste.

11–7. Healthcare facility wastes (general, infectious, pathological, hazardous, and radiological)
   a. Objective. The DA objective is to manage healthcare facility wastes in a manner that protects health and the environment and ensures compliance with appropriate Federal, State, and DA regulations.

   b. Functions. The healthcare facility waste management functions are normally delegated as follows:
      (1) PVNTMED personnel monitor the identification, segregation, transportation, treatment, storage, and disposal of infectious and pathological wastes.
      (2) Healthcare facility personnel ensure the proper handling, identification, segregation, transport, and treatment of infectious, pathological, and general wastes to prevent the potential dissemination of microorganisms. USAEHA TG 147 contains guidelines for use in training personnel in safe handling and disposal of infectious waste.

   c. Requirements.
      (1) General waste.
         (a) Areas generating general waste will follow the same reasonable and prudent methods that are used for collecting refuse generated in nonmedical facilities.
         (b) The waste will be collected at regular intervals by MTF or contractor personnel trained in proper collection and handling procedures.
         (c) Healthcare facility waste handlers should receive initial training in the proper handling and disposal of all wastes. Handlers should receive periodic (at least annual) refresher training on program changes and new developments.
      (2) Infectious waste.
         (a) Units and activities that generate infectious waste will store the waste in the area of generation until collected. Containers with lids (as appropriate) and lined with impervious, tear-resistant, and distinctively colored (for example, red) plastic bags will be used in the area of generation.
         (b) Container liners will be tightly sealed; for example, with twist ties, rubber bands, and/or tape, before leaving the area of generation.
         (c) The waste will be collected at regular intervals by MTF or contractor personnel trained in proper collection and handling procedures.
      (3) Pathological waste.
         (a) Areas that generate pathological waste will handle the waste as outlined in (2)(a) through (c) above.
         (b) When storage of pathological waste is necessary, the enclosed waste will be placed under refrigeration until it is transferred for treatment.
         (4) Transportation within the healthcare facility.
            (a) General waste.
            1. General waste will be transported within the healthcare facility as described for infectious waste in (b) below, unless the facility has a gravity chute or pneumatic tube system. In this case the sealed bags of waste will be placed in a larger plastic bag, sealed, and transported to the system openings.
            2. All general wastes will be taken to outside collection points for subsequent incineration or disposal at an approved sanitary landfill.

            (b) Infectious waste.
            1. Infectious waste will be transported in the original container or the sealed bags can be transferred to larger carts.
            2. Infectious waste will be manually transported to the incinerator or autoclave for treatment.
            3. Carts used to transport infectious waste will be of a type that can be easily cleaned and that will not disclose their contents to hospital personnel. If soiled, containers and carts will be cleaned after each use.
            4. Routes used for transporting infectious waste within the MTF will be carefully selected to minimize patient and personnel exposure and congestion.
            (c) Pathological waste. Pathological waste will be transported and handled the same as infectious waste in (b) above.

      (5) Transportation outside the healthcare facility.
         (a) All Federal, State and local regulations and requirements for transportation manifesting and/or tracking will be followed.
         (b) The transportation of medical wastes in privately owned vehicles is prohibited.

      (6) Treatment and disposal.
         (a) General waste. General waste will be stored outside the healthcare facility in leakproof containers such as dumpsters and compactors.
            1. Containers for storage, transportation, and disposal will be cleaned and sanitized as needed.
            2. The containers (dumpsters) will be transported to the sanitary landfill and must be cleaned before return to the healthcare facility.

         (b) Infectious waste.
            1. Infectious waste will be incinerated or autoclaved (steam) as a method of treatment to render the waste noninfectious.
            2. The ash, or noninfectious waste from treatment by steam sterilization, can be disposed of at the sanitary landfill.
            3. All infectious waste incinerators must be permitted as required by State or local air pollution regulations.
            4. Infectious waste transported to a contract or other off-post incinerator (or other treatment site) will be handled by a manifest procedure whereby a given identifiable quantity of waste is receipted at the disposal site. These receipts are to be returned and made part of the generators’ record of waste production and disposal.

         (c) Pathological waste.
1. Pathological waste will be incinerated or steam sterilized as a method of treatment.

2. After incineration the ash can be disposed of at the sanitary landfill.

3. If steam sterilized, the pathological waste will be subjected to destruction (grinder) and flushed into the sanitary sewer or incinerator.

4. All pathological waste incinerators must be permitted as required by State or local air pollution regulations.

5. Pathological waste transported to a contract or other off-post incinerator (or other treatment site) will be handled by a manifest procedure whereby a given identifiable quantity of waste is receipt accepted at the disposal site. These receipts are to be returned and made part of the generators’ record of waste production and disposal.

(d) Liquid waste. Liquid waste (for example, feces, urine, vomitus, and blood) will be disposed of in the sanitary sewer. Care should be taken to ensure that contamination of hospital personnel and the immediate environment does not occur during disposal. However, liquids from the microbiology laboratory will be steam sterilized before disposal in the sanitary sewer. Liquids from the surgical suite may require steam sterilization at the discretion of the infection control committee before disposal in the sanitary sewer or incinerator.

(e) Needles, syringes, and sharps.

1. Immediately following use, needles, syringes, and other sharps should be placed into rigid, impervious (for example, plastic) containers. These items should not be recapped, clipped, or otherwise intentionally broken or destroyed prior to being placed into these containers.

2. The rigid sharps containers will be clearly labelled to indicate they contain infectious waste consisting of sharps. These containers will be designed to prevent removal of the contents and will be safeguarded by some method (for example, locked in place or under supervision) to prevent misuse or access by unauthorized persons.

3. Sharps containers, when full, will be sealed and transported either separately or in larger infectious waste containers along routes used for other infectious waste. These items will be transported manually to the incinerator, autoclave, or other treatment system. Sharps transported individually to an off-post incinerator (or other treatment site) will be processed by a manifest procedure as described for infectious waste.

4. The treatment system used must render the sharps containers and their contents noninfectious. If, following treatment, the contents of the containers remain intact (for example, potentially reusable), safeguards must exist to prevent reuse of these items while they await final disposal.

d. Contingency plans. Contingency plans will be developed to ensure safe storage, transportation, and disposal in the event the primary method is temporarily disabled or unavailable. This may include written agreements with local civilian hospitals.

11–8. Environmental noise

a. Objective. The DA objective is to control noise from Army sources in a manner that protects community health and welfare without impairing mission or readiness. Specific environmental noise criteria for continuous noise sources are contained in TM 5–803–2/AFM 19–10/NAVFAC P–970. Army policy on land use planning where high noise areas adjacent to Army installations is contained in AR 200–1 and AR 210–20.

b. Functions. Installation PVNTMED personnel will—

(1) Monitor Army operations and activities to ensure compliance with Army and regulatory agency standards.

(2) Provide technical assistance concerning noise abatement, procedures, and controls.

11–9. Spill control

a. Objective. The objective is to prevent the discharge of oil, fuels, and hazardous substances to the environment, and to promptly contain and neutralize such spills.

b. Functions. Installation PVNTMED personnel will—

(1) Review the status of control measures to ensure compliance with health aspects of regulatory agency guidelines.

(2) Provide technical assistance concerning control, containment, and neutralization as appropriate.

(3) Assist installations in preparing spill prevention control and countermeasure plans and installation spill contingency plans.

11–10. Pesticide monitoring

a. Objective. The DA pesticide monitoring program objective is to promote the judicious use of pesticides and ensure the use and disposition of pesticides in a safe manner with minimal health or environmental effect.

b. Functions. USAEHA will—

(1) Investigate all alleged hazardous incidents resulting from the use or disposition of pesticides. As required, USAEHA will conduct scheduled repetitive environmental sampling and analysis for pesticides.

(2) Conduct pesticide monitoring activities that support the national pesticide monitoring program.

(3) Conduct and report special investigations of alleged incidents so that the reports are compatible with the National Environmental Pesticide Data System.

(4) Coordinate the statistical and chemical technologies routinely employed with QA programs of the National Pesticide Monitoring Program.

(5) Periodically evaluate Army pesticide monitoring data to determine common factors that will help implement improved procedures.

Chapter 12 Sanitation

12–1. General

a. Sanitation is one of the most cost-effective means available to the military for preventing disease and improving the soldier’s well-being.

b. This chapter outlines the basic requirements of the Army Sanitation Program. Certain subjects, however, because of specific requirements, are covered elsewhere under separate chapters within this regulation or in TB MEDs, TMs, and FMs referenced and adopted as Army policy by this regulation.

12–2. Fixed installation drinking water program

a. Importance. Few areas of PVNTMED responsibility can have as far reaching an effect as the sanitary control and surveillance of drinking water supplies. In order to ensure the sanitary control of this vital commodity, continuous and effective communication must exist between the facilities engineer and PVNTMED personnel.

b. Fixed installation supplies. Fixed installation supplies are water systems that are enclosed or are protected distribution systems that transfer water from production points to consumption. Sanitary control of fixed installation supplies will be under TB MED 576.

c. Water quality surveillance. Water quality surveillance will be under AR 420–46 and TB MED 576, which includes discussion about compliance with the Safe Drinking Water Act Amendments of 1986, Public Law 99–339 as amended. Installations and/or activities in regions where the Safe Drinking Water Act is applicable will also comply with all subsequent amendments to the act as defined by AR 420–46 and TB MED 576.

(1) In States and territories not having primacy and in States where primary enforcement has been granted by the EPA, Army installations classified as suppliers of water will comply with substantive and procedural requirements of the National Primary Drinking Water Regulation (NPDWR) (AR 420–46 and AR 200–1).

(2) In the OCONUS areas outside those defined in AR 420–46, Army installations classified as suppliers of water will comply with the standards in NPDWR or the host country, whichever are more stringent. Any requests for deviation from the CONUS drinking
water standards will be submitted in writing to the theater surgeon with a copy furnished to HQDA (SGPS–PSP), 5109 Leesburg Pike, Falls Church, VA 22041–3258.


e. Fluoridation. Fluoridation of water will be according to AR 420–46 and TB MED 576.

f. Cross connections. Cross connections between potable and nonpotable distribution systems are not permitted. TB MED 576 and TM 5–660/AFR 91–26/NAVFAC MO–210 discuss cross connections and provide proper references. The current National Standard Plumbing Code will be followed in the design, maintenance, and renovation of water distribution systems and in the selection of all plumbing fixtures.

g. Installation commander functions. This commander ensures that appropriate samples are collected from all potable water sources and are analyzed for chemical parameters by a certified laboratory in CONUS and validated by USAEHA’s laboratory QA program in OCONUS laboratories. All test data are forwarded to appropriate regulatory authorities. Copies are provided to the IMA for medical review and/or evaluation and to the Commander, USAEHA, AT- TN, HSHB–ME–W, Aberdeen Proving Ground, MD 21010–5422, for additional evaluation and for inclusion in the Army’s drinking water surveillance program data base. USAEHA can provide technical and/or operational assistance for such requirements. The USAEHA operates a laboratory, certified by Federal and State regulatory authorities, to perform analyses of specially requested services. Details of these activities are in TB MED 576.

h. PVNTMED functions. The PVNTMED will—

(1) Provide medical evaluation of monitoring data for the potable water supply and distribution system as necessary to fulfill the requirements of NPDWR, National Secondary Drinking Water Regulations (as applicable), and TB MED 576. USAEHA can provide technical assistance for such activities, and maintains a laboratory certified by Federal and State regulatory authorities to support specially requested services. (Routine analyses obtained to comply with regulatory requirements will be the responsibility of the installation commander and engineers. The responsible IMA and USAEHA will receive copies of all potable water analyses, because engineers must coordinate with and obtain necessary assistance from the PVNTMED on all matters pertaining to health and esthetic aspects of regulatory compliance. This process is detailed in TB MED 576.)

(2) In cooperation with the engineers, maintain liaison with proper Federal, State, and local regulatory authorities regarding current drinking water regulations.

(3) Interpret results of water quality analyses.

(4) Approve concentrations and types of chemical additions to potable water supplies.

(5) Maintain records, under NPDWR and AR 25–400–2, that reflect the chemical, radiological, and microbiological quality of the installation potable water.

(6) Conduct programmed sanitary inspections of the entire potable water system on a yearly basis.

(7) Perform special sanitary surveys as conditions warrant.

(8) Develop information whereby installation personnel and family members can be notified of any degradation or contamination of the potable water system and recommend appropriate corrective action to the installation commander.

(9) Perform independent surveillance of Government-owned, contractor-operated facilities per TB MED 576.

(10) Conduct bacteriological, concurrent chlorine residual, and fluoride surveillance analysis (if applicable) of the potable water system for supplied and purchased sources under the requirements of NPDWR, AR 420–46, and TB MED 576.

(11) Provide information and guidance to the installation commander concerning the following:

(a) Current requirements for, availability of, and regulations concerning potable water.

(b) The need for and method of water conservation.

(c) Available methods to reduce pollution of water supply by installation activities.

12–3. Ice manufacture

a. In CONUS, Alaska, and Hawaii, sanitary inspections of ice manufacturing facilities are the responsibility of the U.S. Army Veterinary Service. When circumstances dictate, and water potability certification by the military is necessary, veterinary personnel will collect samples and submit them to the appropriate Army PVNTMED activity for testing. Results of the tests will be provided to veterinary personnel for use in completing their inspection report.

b. In OCONUS, PVNTMED personnel will conduct sanitary inspections of ice manufacturing, storage, and distribution facilities; and will, in coordination with appropriate veterinary personnel, recommend approval of commercially-operated plants. Approved commercial ice plants will be listed in the Directory of Sanitarily Approved Food Establishments for Armed Forces Procurement or in the locally approved establishment list (AR 40–657/NAVSUPINST 4355.4/AFR 161–32/MCO P10110.31).

c. Sanitary requirements for ice manufacture are detailed in MIL STD 906. Additional requirements follow:

(1) Plumbing installation will be according to the requirements of the current National Standard Plumbing Code or local jurisdiction plumbing code, whichever is more strict.

(2) Surfaces of floors, walls, and ceilings of all rooms used for manufacture, processing, and storage of ice will be smooth, impervious, and nontoxic (under use conditions).

(3) All can fillers, coresucking devices, and drop tubes will be handled in a manner to prevent contamination. Freezing cans will be disinfected by steam or by being submerged for 2 minutes in a 100 parts per million (ppm) free available chlorine solution.

(4) Only dedicated vehicles will be used for transporting ice. An exception is granted for transportation of packaged and/or containerized ice in enclosed clean multi-use vehicles.

12–4. Water supply afloat

All Army floating vessels will use the procedures contained in AR 56–9 to provide and maintain a safe and sanitary water supply. Bacteriological, physical, and chemical requirements for potable water will meet the criteria in AR 56–9 and TB MED 576. Assistance in monitoring these requirements will be provided by the MEDDAC and/or MEDCEN PVNTMED service supporting the vessel’s home port or port of call.

12–5. Field water supply program

a. Significance. Field water supplies refer to water systems of a nonpermanent nature used in training exercises and actual operations. The water may be transported from point of production to point of use in temporary conduits or unprotected portable containers. The provision of and sanitary control of potable water in the field is of significant importance to the well-being and morale of all concerned.

b. Sanitary control. Sanitary control of field water supplies will be according to AR 700–136, FM 10–52, FM 21–10/AFM 161–10, and TB MED 577.

12–6. Swimming pools

a. Guidance. Sanitary control and operation of Army swimming pools and swimming areas will be under AR 420–46, TM 5–662, and TB MED 575.

b. PVNTMED functions. Installation PVNTMED personnel will—

(1) Assist the installation commander in the sanitary control of swimming facilities.

(2) Maintain current information that includes engineering plans, type, location, size, maximum bather load, and operating hours for all swimming facilities.

(3) Provide training for lifeguards and applicable facilities engineer personnel in the sanitary operation and monitoring of swimming facilities. PVNTMED personnel also will ensure that all

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of air-conditioning, evaporative cooling, dehumidification, and mechanical ventilation equipment and systems are contained in DOD 4270.1–M and TM 5–810–1.

12–8. Barber and beauty shops
Sanitary requirements for barber and beauty shops are detailed in appendix D.

12–9. Drycleaning
General guidance related to customer-operated and commercial dry-cleaning operations can be obtained from the Commander, USAEHA, Aberdeen Proving Ground, MD 21010–5422.

12–10. Mobile home parks
Sanitary requirements for mobile home parks are described in appendix E.

12–11. Child development services facilities
Guidance relative to the sanitary requirements for CDS facilities is in AR 608–10. Environmental sanitation inspections will be conducted monthly for center-based CDS facilities. For quarters-based family child care homes, see AR 608–10.

12–12. Recreational areas
a. Recreational areas must be constructed and operated to provide a pleasant, safe, and sanitary recreational area while at the same time providing adequate protection of the environment.

b. A comprehensive presite selection of recreational areas is required. General requirements for site selection and development are outlined in Public Health Service Publication No. 1195 and TM 5–803–12. Particular attention must be placed on providing an adequate potable water supply, waste disposal, drainage, prevention of soil erosion, and protection of watersheds.

c. Natural bathing areas and swimming pools will be designed and operated under AR 420–46, TB MED 575, and TM 5–662.

d. Potable water will be supplied to all recreational areas except those designated as wilderness areas.

e. Adequate solid waste disposal will be provided. Containers will be emptied and cleaned on a schedule approved by the IMA.

f. Liquid waste disposal will be through a sanitary sewer.

g. Nonwaterborne waste disposal systems will not be used unless approved by the IMA. When approved, nonwaterborne waste disposal facilities will be vermin-proofed, equipped with self-closing doors, adequately screened, and protected from inclement weather. Handwashing facilities should also be provided. The IMA will participate in establishing the cleaning and emptying schedules (as appropriate) with installation facilities engineer personnel.

12–13. Laundry operations
a. Fixed laundry operations will be designed and operated under AR 420–130.

b. Field laundry operations will be designed and operated under FM 10–280.

12–14. Sports facility sanitation
a. Environmental considerations concerning Army sports facilities include prevention of infections due to contamination of equipment, towels, clothing, and other common use items, and provision of adequate facilities and housekeeping.

b. Common use items such as athletic shoes must be disinfected with an approved fungicide spray and must be air dried thoroughly before being reissued. Towels and issued athletic clothing must be laundered before being reissued.

c. Whirlpool baths, steam cabinets, and other therapy-type equipment must be disinfected between users using either a disinfectant solution containing a minimum of 50 ppm free available chlorine or an iodine disinfectant providing the equivalent of 25 ppm available iodine. Equipment must be rinsed with potable water after disinfection.

d. Showers and locker room floors and benches will be cleaned and disinfected at least daily. Toilet facilities will be cleaned at least
daily. Disinfectant products will be applied according to manufacturers’ instructions.

- Athletic fields will be provided with adequate potable water supplies and convenient latrine facilities.
- Temporary and mobile food services will be operated according to paragraph 12–17 of this regulation and TB MED 530.

12–15. Confinement facilities

- a. Sanitary inspection requirements for Army detention and confinement facilities are outlined in AR 190–38 and AR 190–47. The IMA will provide required support to carry out the requirements outlined in these ARs.
- b. Particular attention must be given to providing adequate floor space, temperature control, ventilation, and housekeeping in detention cells, isolation rooms, prisoner dormitories, dining facilities, and other common use areas.

12–16. Disinfectant selection

- a. Complete evaluation of disinfectant products for microbiological effectiveness is beyond the capability of most PVNTMED services and should not be undertaken. Disinfectants, sanitizers, and other chemical or physical agents designed to reduce or inhibit the growth of microorganisms may be used in food service facilities, hospitals, and other areas as appropriate. Only products that have been approved by the EPA, the Food and Drug Administration (FDA), the USDA, or any combination of the above three agencies as required by Federal law will be used.
- b. Selection of a disinfectant, sanitizer, or other such product is dependent on the following criteria:
   - (1) The product is approved for the intended use.
   - (2) The product is compatible with the local water supply considering hardness, hydrogen ion concentration (pH), and other physical and chemical parameters.
   - (3) The product is safe, nonirritating, and nontoxic when used according to directions.
   - (4) The product must perform the intended and stated task satisfactorily.

12–17. Food service sanitation

- a. The attainment of quality food service is paramount at all levels of command. Food is easily contaminated and will readily support the growth of many disease-producing microorganisms. The trend to larger facilities and centralized preparation increases the potential for large-scale illness and resultant loss of mission effectiveness.
- b. The essential elements to be implemented for the food service sanitation program are in TB MED 530. Additional guidance on expedient methods for field food service operations is presented in FM 8–250 and FM 21–10/AFM 161–10.
- c. Veterinary personnel will conduct necessary sanitary inspections described in AR 40–657/NAVSUPINST 4355.4/AFR 161–32/MCO P10110.31 and other inspections related to veterinary aspects of procurement, processing, storage, shipment, receipt, and distribution of food. PVNTMED personnel will assess the adequacy of food service sanitation practices and storage of food products at the food preparation facility.

12–18. Commissary and troop issue subsistence activity sanitation

Commissary and troop issue subsistence activity sanitation procedures and policies are presented in the Commissary Operations Manuals, MIL STD 668, MIL STD 903, and in TB MED 530. Veterinary personnel will conduct the necessary sanitation inspections relating to the procurement, processing, storage, shipment, receipt, and distribution of food.

Chapter 13
Environmental Laboratory Services

13–1. General

- a. Quality analytical chemistry laboratory support is needed to assess the nature and extent of potential hazards in the environment of personnel served by the PVNTMED service of each MEDDAC or MEDCEN. This need is reflected in local, State, and Federal requirements that apply to environmental chemistry laboratory support.
- b. This chapter gives guidance to PVNTMED services for effective laboratory support to fulfill their operational requirements to promote health and prevent disease.

13–2. Coordination

- a. The resources of the local PVNTMED service are the first to be considered by a MEDDAC or MEDCEN in dealing with an environmental problem.
- b. If local PVNTMED service resources are not adequate to meet requirements, the USAEHA or the OCONUS support laboratory will provide support per paragraph 1–8.
- c. Coordination with USAEHA or the OCONUS supporting laboratory will be made before sample collection unless the samples are to be analyzed locally.

13–3. PVNTMED service

- a. Laboratory services provided by PVNTMED services in each MEDDAC and/or MEDCEN will vary according to the size and location of the installation, mission of the units supported, and personnel assigned. The local service will establish, as a minimum, a laboratory capability to monitor fluoride, chlorine residual, pH, and bacteriological parameters to ensure adequate surveillance and sanitary control of installation water supplies and recreational waters (TB MEDs 575 and 576).
- b. Surveillance of chemical, pesticide, and radiological levels in drinking waters will be performed per TB MED 576.

13–4. Certification and/or accreditation

- a. Federal facilities within the 50 States, the District of Columbia, and those OCONUS areas specified in AR 420–46 must comply with all Federal, State, and local requirements regarding the Safe Drinking Water Act Amendment of 1986 (Public Law 99–339), as implemented by EPA in 40 CFR 141 and 142. For the purposes of certification, Federal laboratories that conduct routine monitoring of public drinking water supplies are to be considered local laboratories and will be certifiable under a State or regional EPA program. Details on the EPA’s water supply laboratory certification program and the technical criteria considered essential to generate valid data are found in EPA 5709–82–002. Commanders of installations located elsewhere will assure compliance with TB MED 576 as interpreted by the local medical authority.
- b. Per AR 420–46, laboratory facilities performing operational (such as microbiological and chemical) analyses will comply with substantive and procedural requirements, if any, issued by State or regional EPA authorities.

13–5. Quality assurance

- a. Data reliability. The role of the environmental laboratory is to provide qualitative and quantitative data to be used in decision-making. Because of the importance of laboratory analyses and the resulting actions they produce, a program to ensure the reliability of the data is essential.
- b. Quality control of laboratory analyses.

  (1) An established, routine, quality control program applied to every analytical test is important to the reliability of the final results. Quality control guidelines for laboratories performing water and wastewater analyses are in EPA handbook 600/4/79–019, produced by the Analytical Quality Control Laboratory, National Environmental Research Center, Cincinnati, Ohio, and in EPA manual 600/8/78–017, produced by the Environmental Monitoring and Support
Laboratory, Cincinnati, Ohio (EMSL–CI). Known value quality control samples for microbiology and chemistry will be furnished by EMSL–CI directly to EPA regions and through the regions to local laboratories. Requests for samples will be made through the appropriate EPA regional office.

(2) USAEHA provides proficiency testing surveys for quality control of water and wastewater analyses. PVNTMED services and OCONUS supporting laboratories wishing to participate in this program should address their requirements to the Commander, USAEHA, ATTN: HSHB–ML–A, Aberdeen Proving Ground, MD 21010–5422.

(3) AR 750–25 establishes a single DA test, measurement, and diagnostic equipment (TMDE) calibration and repair support program to ensure that all TMDE receive calibration on a recurring basis as required. TB 750–25 describes procedures and responsibilities for obtaining, providing, and receiving calibration service.

Chapter 14
Field Preventive Medicine

14–1. General

a. Preventable personnel losses from heat, cold, or disease become important because history has repeatedly shown that nonbattle losses have played a significant role in the outcome of military operations. This chapter establishes field PVNTMED responsibilities.

b. Guidance herein applies to field training exercises, disaster relief operations, as well as contingency force deployment. Included are—

(1) Individual preventive medicine measures (PMM).

(2) Company-level PMM.

(3) Division-level PMM.

(4) PVNTMED team support.

14–2. Individual support

The soldier will employ all protective measures possible. Each soldier, as a minimum, will protect against—

a. Heat incapacitation in hot climates by drinking a sufficient volume of water at frequent intervals.

b. Cold injury in cold climates by wearing proper cold-weather clothing and frequently changing socks to keep feet dry, by careful handling of gasoline-type liquids, and by avoiding contact between skin and cold metal.

c. Mosquito, fly, tick, and other arthropod-borne diseases by using insect repellents, netting, and insecticide aerosols; by taking approved chemoprophylaxis; and by wearing the uniform properly.

d. Enteric disease by using iodine tablets whenever water quality is uncertain, by avoiding unapproved civilian food vendors, and by properly disposing of bodily wastes.

e. Skin disease by washing the body as often as practicable.

f. Other hazards by using appropriate measures as described in FM 21–10/AFM 161–10.

14–3. Company-level PMM

Companies, troops, batteries, and units of equivalent size are responsible for those PMM that affect units as a whole or are beyond the resources of an individual soldier. FM 21–10/AFM 161–10 will be used as a guide with maximum use of company-level PMM therein. Commanders will ensure that their units conduct PMM.

a. Functions. As a minimum, units deploying to the field will—

(1) Before deployment, appoint a field sanitation team with responsibilities defined in b below.

(2) Before deployment, incorporate PMM into SOPs.

(3) Have the capability to use pesticides and vegetation controls.

(4) Bury and/or burn wastes to prevent the breeding of insects and rodents. Consult the environmental coordinator or PVNTMED personnel to ensure compliance with local environmental regulations and laws during field exercises.

(5) Protect food during storage and preparation to prevent contamination (TB MED 530).

(6) Monitor unit water sources to assure adequate supplies and disinfection.

(7) Arrange for maintenance of immunizations and prophylaxis.

(8) Use other appropriate measures under FM 21–10/AFM 161–10.

(9) Assure command supervision of individual PMM.

(10) Request assistance for problems exceeding unit capabilities.

(11) Deploy to the field with field sanitation equipment listed in table 14–1.

b. Field sanitation teams.

(1) When organic or attached medical personnel are available, they will be appointed and will serve as the field sanitation team for the unit. They will serve as advisers to the commander, train unit personnel in individual PMM, and supervise or conduct basic PVNTMED services. Company and battery-sized units deploying without organic or attached medical personnel will appoint a field sanitation team. These field sanitation teams provide the small-unit commander with—

(a) Organic expertise to monitor the status of unit PMM.

(b) Limited capability to control insect and rodent vectors in the unit area.

(2) The field sanitation team will conduct training within the unit on individual PMM against disease and injury as these relate to an assessment of the medical threat in the prospective or defined areas of operation.

(a) Composition. Company aidmen (military occupational specialty 91A) organic or attached to deployed units will be trained and will function as the unit field sanitation team. If medical personnel are not available, two soldiers will be selected and trained, one of whom must be a noncommissioned officer. Neither the organic or attached aidman nor selected soldiers will have less than 6 months remaining with the unit on the date of appointment.

(b) Training. Members of field sanitation teams (organic or attached medical or nonmedical personnel) will receive training from supporting medical resources before deployment or field exercises to assure that small units have the PVNTMED resources to operate in adverse disease and/or climatic environments. Instruction will address use, maintenance, and care of the field sanitation team equipment as well as communicable disease control, food service sanitation, water supply, waste disposal, and arthropod and rodent control.

(c) Coordination. In nonoperational areas, pesticide spraying is generally accomplished by facilities engineer personnel. Such services will reduce the field sanitation team workload. Teams should coordinate with engineer personnel. Field sanitation team spraying will be conducted as necessary to supplement engineer coverage and to maintain team familiarity with control techniques. Pesticide recommendations to control specific vectors or pests are found in AFPMB TIM 24. Large populations of insect or rodent pests in the unit area will warrant a request for support by a PVNTMED (LA) team. (See para 14–5d(2) below.)

(d) Equipment. The recommended stocking of field sanitation team materials is listed in table 14–1. All deployable units will maintain, transport, and use listed items in support of both training exercises and contingency mission.

14–4. Division-level PMM

The next echelon of PVNTMED support is at the division. This level of support is provided by the PVNTMED section in the medical battalion of the infantry (light), airborne, and air assault divisions. In armored and mechanized divisions, this support is provided by PVNTMED elements of the main support battalion. The services provided include identification of PVNTMED problems and training of unit field sanitation teams and back-up company-level PMM which are beyond the capability of unit personnel due to their complexity, scope, or specialized nature.

14–5. PVNTMED team support

Although the main thrust of PVNTMED occurs within small units
in the form of individual and company-level PMM, some problems will require additional expertise and equipment for resolution. Such skills and materiel are concentrated in PVNTMED teams that can be used to support units operating in areas of highest disease risk.

Table 14–1
Field sanitation team materials

| National stock number: 6810–00–255–0471 | Description: Calcium Hypochlorite, 6 oz | Unit/Issue: BT | Allowance: 3 | Authority: CTA 50–970 |
| National stock number: 6545–00–914–3480 | Description: Chest, No. 3, 30x18x10, Alum | Unit/Issue: EA | Allowance: 1 | Authority: CTA 8–100 |
| National stock number: 6850–00–270–6225 | Description: Chlorination Kit, Water purification | Unit/Issue: KT | Allowance: 10 | Authority: CTA 50–970 |
| National stock number: 6840–00–810–6396 | Description: Disinfectant, food service, 12’s | Unit/Issue: BX | Unit/Issue: 2 | Authority: CTA 50–970 |
| National stock number: 3740–00–132–5936 | Description: Duster, Manually operated, tubular pump | Unit/Issue: EA | Allowance: 1 | Authority: CTA 50–970 |
| National stock number: 8415–01–012–9294 | Description: Glove, Chemical and Oil Protective | Unit/Issue: PR | Allowance: 1 | Authority: CTA 50–900 |
| National stock number: 4240–00–190–6432 | Description: Goggles, Industrial, non-vented | Unit/Issue: PR | Allowance: 384 | Authority: CTA 50–970 |
| National stock number: 6840–01–284–3982 | Description: Insect Repellent, personnel application, 2 oz | Unit/Issue: BT | Allowance: 384 | Authority: CTA 50–970 |
| National stock number: 6840–01–210–3392 | Description: Insecticides, Chlorpyrifos, 42%, 40 ml, 12’s | Unit/Issue: EA | Allowance: 1 | Authority: CTA 50–970 |
| National stock number: 6840–01–067–6674 | Description: Insecticides, d-Phenothrin, 2%, 12 oz | Unit/Issue: CN | Allowance: 144 | Authority: CTA 50–970 |
| National stock number: 6840–00–242–4217 | Description: Insecticides, lindane, 1%, 2 oz | Unit/Issue: BT | Allowance: 192 | Authority: CTA 50–970 |
| National stock number: 3740–00–252–3384 | Description: Mousetrap, spring, 12’s | Unit/Issue: DZ | Allowance: 2 | Authority: CTA 50–970 |
| National stock number: 3740–00–260–1398 | Description: Rattrap, spring, 12’s | Unit/Issue: EA | Allowance: 2 | Authority: CTA 50–970 |

Table 14–1
Field sanitation team materials—Continued

| National stock number: 3740–01–234–3448 | Description: Rodenticides, anticoagulant bait, 5 lb | Unit/Issue: CN | Allowance: 2 | Authority: CTA 50–970 |
| National stock number: 6840–00–753–4973 | Description: Swatter, fly, 12’s | Unit/Issue: PG | Allowance: 1 | Authority: CTA 50–970 |
| National stock number: 6850–00–985–7166 | Description: Water purification tablet, iodine, 50’s | Unit/Issue: BT | Allowance: 400 | Authority: CTA 8–100 |

Notes:
1 Unit/Issue entries are computer entry codes; for example, BT is the code for bottle and PG is package.

a. PVNTMED teams will—
(1) Prepare for deployment in support of contingency or disaster relief operations within 24 hours after notification. Predeployment measures will include—
(a) Readiness of authorized equipment and vehicles.
(b) Stocking of prescribed expendables.
(c) Procurement of CTA 50–900 equipment for augmentation personnel.
(d) Rehearsal of loading plans.
(e) Coordination with the Armed Forces Medical Intelligence Center and Armed Forces Pest Management Information Analysis Center, as applicable, concerning epidemiologic and vector information on regions of likely deployment.

(2) Participate in field training exercises, as directed. Staff coordinating procedures will be emphasized to assure that the supported unit will make full use of technical resources if strategic deployment becomes necessary.

(3) Not normally be used as field sanitation teams for hospitals, headquarters, or other units to which attached.

b. HSC will—
(1) Maintain a roster of professional fillers to facilitate augmentation of Forces Command PVNTMED teams in the event of disaster relief operations or contingency force deployment.

(2) Schedule annual liaison visits for key filler personnel to facilitate rapport and to assure both individual and team deployment readiness.

c. USAPACEHEA will maintain an active capability to—
(1) React to requests from Commander in Chief, U.S. Army, Pacific (CINCUSARPAC) for assistance in disaster relief operations. Perform necessary liaison with key CINCUSARPAC personnel to assure effective disaster relief response.

(2) Perform basic field PVNTMED operations in areas of mobilizing troop concentrations (within the Pacific Theater); until PVNTMED teams organic to the mobilizing force can arrive on site.
and become mission effective. Participate in field exercises to ensure this capability is well practiced.

d. The full range of PVNTMED services will be provided.
   (1) Command and control, AM teams (if deployed). The AM teams will provide command and control for PVNTMED teams in the area of operations. The operations of individual teams will be decentralized, whenever possible, with attachment to corps elements at greatest risk.

   (2) Entomology service, LA teams. The LA teams will provide entomology support for all elements in the area of operations with support priority to combat units. During the strategic deployment phase of contingency operations, emphasis will be on preemptive suppression of high risk vectors near assembly areas and along routes of movement. As tactical elements deploy for action, LA teams will be prepared to extend services in response to requests from division and corps surgeons.

   (3) Environmental sanitation service, LB teams. The division preventive medicine section will monitor the status of company-level PMM. They will provide consultative support as necessary, to include the onsite training of field sanitation teams. Support priority will go to combat units. During the strategic deployment phase of contingency operations, emphasis will be on assembly areas near points of debarkation. As tactical elements deploy for action, LB teams will be prepared to shift emphasis forward to brigade and/or battalion trains in response to requests from division surgeons. LB teams will coordinate with civil affairs and military police to assure that refugee enclaves and prisoner compounds do not become foci of epidemic disease.

   (4) Environmental engineering service, LC teams. The LC teams will provide the same services cited for LB teams. During the strategic deployment phase of contingency operations, emphasis will be on points of debarkation (airfields and ports). As corps elements deploy from initial staging areas, support priority will shift to corps support groups and area support groups as they become established along lines of communication. LC teams will coordinate with LB teams to assure optimum use of technical resources in the corps as a whole. LC teams will coordinate with the engineers and civil affairs to assure restoration of water, wastewater treatment, and waste disposal facilities in towns and villages within the area of operations.

   (5) Epidemiology service, LD teams. The LD teams will provide epidemiologic services in the area of operations. During the strategic deployment phase of contingency operations, emphasis will be on population centers near assembly areas and along proposed routes of movement to assure early detection of disease. As corps elements deploy for action, support priority will shift to combat units in response to requests from division surgeons.

   (6) Entomology laboratory, LE teams. The LE teams will provide entomological laboratory support for LA teams in the area of operations with initial emphasis on pesticide resistance assessments to facilitate early procurement of alternative pesticides, if warranted.

   (7) Veterinary, JA/JB teams. The JA/JB teams will provide veterinary medical support in areas of operations with initial emphasis on wholesomeness and acceptability of food supplies or sources.

      (a) Initial emphasis will concentrate on protection of the disrupted populace from potentially epidemic diseases.

      (b) Early coordination with host nation public health authorities will be accomplished to assure consonance with local environmental constraints and to permit use of local resources.

      (c) Overall emphasis will be on restoration of local capacity for self-support.
References

Section I
Required Publications

AMH (JCAHO)
Accreditation Manual for Hospitals, Joint Commission on Accreditation of Healthcare Organizations. (Cited in para 8–4b.) (May be obtained from the JCAHO, 875 N. Michigan Avenue, Chicago, IL 60611.)

ANSI Z87.1
Practice for Occupational and Educational Eye and Face Protection. (Cited in paras 5–4l and 5–15c.) (May be obtained from the American National Standards Institute Incorporated, 1430 Broadway, New York, NY 10018.)

AR 10–5
Department of the Army. (Cited in paras 3–4d and 3–10.)

AR 11–34
The Army Respiratory Protection Program. (Cited in para 5–26c(2)(c)).

AR 25–400–2
The Modern Army Recordkeeping System (MARKS). (Cited in paras 1–7,4–5h(17), 5–16b(2)(c),5–21a and c, 9–7a(4) and (6), 9–9a(2)(c) and b(8), and 12–2h(5).)

AR 40–12/SEACNAVINST 6210.2/AFR 161–4
Medical and Agricultural Foreign and Domestic Quarantine Regulations for Vessels, Aircraft, and Other Transports of the Armed Forces. (Cited in para 10–16.)

AR 40–14/DLAR 1000.28
Control and Recording Procedures for Exposure to Ionizing Radiation and Radioactive Materials. (Cited in paras 9–3a,9–4a and b; 9–5a; 9–6a; and 9–9a(2)(a) and (b) and b(1) and (6).)

AR 40–35
Preventive Dentistry. (Cited in para 1–4h(1).)

AR 40–46
Control of Health Hazards from Lasers and Other High Intensity Optical Sources. (Cited in paras 9–4b(1); 9–5b; and 9–9a(2)(a), (b), and (f)).

AR 40–63/AFR 167–3
Ophthalmic Services. (Cited in para 5–15b.)

AR 40–66
Medical Record and Quality Assurance Administration. (Cited in paras 4–11b; 5–16b(2)(c);and 5–21a, d(1), and f.)

AR 40–68
Quality Assurance Administration. (Cited in para 4–9e.)

AR 40–400
Patient Administration. (Cited in paras 3–1; 3–2;3–12a(4); 4–5b(4), c(4), and d(3); 6–3c(7); 9–9c(8);9–12b(3)(a); B–2c; and B–3b.)

AR 40–407
Nursing Records and Reports. (Cited in paras 3–15 and 6–5a.)

AR 40–501
Standards of Medical Fitness. (Cited in paras 5–9b, 5–20b(3) and (5),5–26c(2)(b), and B–4.)

AR 40–562/NAVMEDCOMINST 6230.3/AFR 161–13/CG COMDTINST M6230.4D
Immunizations and Chemoprophylaxis. (Cited in paras 4–3a(3); 4–4,4–5a(3)(c), b(3), d(2)(b), and f(2)(a); and 5–12.)

AR 40–574/AFR 91–22
Aerial Dispersal of Pesticides. (Cited in para 10–12.)

AR 40–657/NAVSUPINST 4355.4/AFR 161–32/MCO P10110.31
Veterinary/Medical Food Inspection and Laboratory Service. (Cited in paras 12–3b and 12–17c.)

AR 50–5
Nuclear Surety. (Cited in para 5–19.)

AR 50–6
Chemical Surety. (Cited in para 5–19.)

AR 55–38/NAVSUPINST 4610.33/AFR 75–18/MCO 4610.19/DLAR 4500.15
Reporting of Transportation Discrepancies in Shipments. (Cited in para 9–9b(13).)

AR 56–9
Watercraft. (Cited in para 12–4.)

AR 95–3
General Provisions, Training, Standardization, and Resource Management. (Cited in para 5–16b(3).)

AR 200–1
Environmental Protection and Enhancement. (Cited in paras 1–6a, 9–3b, 10–4d,10–4(i),11–2b(3), 11–8a, and 12–2c(1).)

AR 210–10
Administration. (Cited in para 4–5e(2)(e).)

AR 210–11
Installations—Billeting Operations. (Cited in para 12–7b(1).)

AR 210–20
Master Planning for Army Installations. (Cited in paras 11–8a and 12–7a.)

AR 210–30
Selection of Sites for Army Installations. (Cited in para 12–7a.)

AR 385–10
Army Safety Program. (Cited in paras 1–4a and c,5–4a(5)(b), b, and f; 5–11;5–15c; 5–16b(1)(a), (4),(5)(b) and (c), (8)(e), and(9)(f); 5–26a; 8–2a;9–7a(2); 9–9a(1); and 10–14.)

AR 385–11
Ionizing Radiation Protection ( Licensing, Control, Transportation, Disposal, and Radiation Safety). (Cited in paras 9–4a and b(1);9–9a(2)(f), and b(2)(3), (12), (13), and (14); and 9–10b and c.)

AR 385–30
Safety Color Code Markings and Signs. (Cited in paras 5–16b(6)(a) and 9–9a(2)(d).)

AR 385–32
Protective Clothing and Equipment. (Cited in para 9–7a(2).)

AR 385–40
Accident Reporting and Records. (Cited in paras 5–4b, 5–10b, 5–22c, 8–2b,9–6a(3), 9–9b(6) and c(8), and 9–12b(3)(b).)

AR 420–10
Management of Installation Directorate of Engineering and Housing and Personnel. (Cited in para 10–3c.)
NBSH 114
General Safety Standards for Installations Using Non-Medical X-Ray and Sealed Gamma-Ray Sources, Energies up to 10 Mev. (Cited in para 9–9(a)(10).)

NFPA Std 70
National Electrical Code. (Cited in para 8–4(b)(1).) (This and the National Fire Prevention Association publication listed below may be obtained from the National Fire Prevention Association, Batterymarch Park, Quincy, MA 02269.)

NFPA Std 99
Standard for Health Care Facilities. (Cited in para 8–4(b)(1).)

NIOSH Publication No. 75–137
Development and Evaluation of Methods for the Elimination of Waste Anesthetic Gases and Vapors in Hospitals. (Cited in para 8–4(b)(5).) (This and the National Institute for Occupational Safety and Health publications listed below may be obtained from the Superintendent of Documents, Government Printing Office, Washington, DC 20402.)

NIOSH Publication No. 77–140
Occupational Exposure to Waste Anesthetic Gases and Vapors. (Cited in para 8–4(b)(5).)

NIOSH Publication No. 77–171
Control of Occupational Exposure to N2O in the Dental Operator. (Cited in para 8–4(b)(5).)

NIOSH Publication No. 77–200
Special Occupational Hazard Review with Control Recommendations—Use of Ethylene Oxide as a Sterilant in Medical Facilities. (Cited in para 8–4(b)(5).)

NIOSH Publication No. 81–123
Occupational Health Guidelines for Chemical Hazards. (Cited in para 5–9(a)(2).)

OSHA 2014
Recordkeeping and Reporting Guidelines for Federal Agencies. (Cited in para 5–22(c).) (May be obtained from the Office of Federal Agency Programs, 200 Constitution Avenue, NW, Washington, DC 20210.)

OTSG Policy Letter 86–01.0
Surveillance of Laser and Microwave/Radiofrequency Workers. (Cited in para 5–15(a)(4).) (May be obtained from the Commander, USAEHA, ATTN: HSHB–MS, Aberdeen Proving Ground, MD 21010–5422.)

Public Health Service Publication No. 1195
Environmental Health Practice in Recreational Areas. (Cited in para 12–12b,(May be obtained from the Interagency Program Retail Food Protection Branch, Food and Drug Administration, 200 C Street, SW, Washington, DC 20204.)

SB 3–40
Pesticides. (Cited in para 10–4.)

TB MED 2
Sterilizing Medical, Surgical, Dental and Veterinary Material. (Cited in para 8–4(b)(4).)

TB MED 3
Occupational Health and Safety in Dental Clinics. (Cited in para 8–4(b)(5).)

TB MED 81
Cold Injury. (Cited in para B–3.)

TB MED 266
Disinfection and Sterilization of Dental Instruments and Materials. (Cited in para 8–4(b)(5).)

TB MED 501
Occupational and Environmental Health: Hearing Conservation. (Cited in paras 5–9(a)(4); 5–16 and (7); 5–26(c)(3); 5–26(b)(2).) (Cited in para 8–4(b)(5).)

TB MED 502
Occupational and Environmental Health: Respiratory Protection Program. (Cited in paras 5–9(a)(5); 5–26(c)(2)(c); 8–4(b)(5); 9–7(a)(1), (3), and (4); and 10–14.)

TB MED 503
The Army Industrial Hygiene Program. (Cited in para 5–4(d)(4); 5–4(h)(4), 16–16(b)(8); 5–24, and 8–4(b)(5).)

TB MED 506
Occupational and Environmental Health: Occupational Vision. (Cited in paras 5–9(a)(6) and 5–15a and a(7).)

TB MED 507
Occupational and Environmental Health: Prevention, Treatment, and Control of Heat Injury. (Cited in para B–2.)

TB MED 509
Spirometry in Occupational Health Surveillance. (Cited in paras 5–9(a)(7) and 5–26(c)(2)(c).)

TB MED 510
Interim Guidelines for the Evaluation and Control of Occupational Exposure to Waste Anesthetic Gases. (Cited in paras 5–9(a)(8) and 8–4(b)(5).)

TB MED 513
Guidelines for the Evaluation and Control of Asbestos Exposure. (Cited in paras 5–4(a)(6), 5–4(e)(5), 5–9(a)(9), and 5–28a and b.)

TB MED 521
Management and Control of Diagnostic X-Ray, Therapeutic X-Ray, and Gamma-Beam Equipment. (Cited in paras 9–8(a)(7) and 9–9(a)(2)(d) and b(7), (8), and (9).)

TB MED 522
Occupational and Environmental Health: Control of Health Hazards from Protective Material Used in Self-Luminous Devices. (Cited in para 9–10c.)

TB MED 523
Control of Hazards to Health from Microwave and Radio Frequency Radiation and Ultrasound. (Cited in paras 5–9(a)(10), 9–5(b), and 9–9(c)(5).)

TB MED 524
Occupational and Environmental Health: Control of Hazards to Health from Laser Radiation. (Cited in paras 5–9(a)(11), 9–5(b), 9–7(b), and 9–9(c)(5).)

TB MED 525
Control of Hazards to Health from Ionization Radiation Used by the Army Medical Department. (Cited in paras 9–4 and b–9–8(a)(6); 9–9(a)(2)(d) and (f), b(2) and (3); and 9–10a.)
TB MED 530  
Occupational and Environmental Health: Food Service Sanitation. (Cited in paras 10–9a, 12–14f, 12–17b, 12–18, and 14–3a(5).)

TB MED 575  
Swimming Pools and Bathing Facilities. (Cited in paras 12–6a and b(5) and (6), 12–12c, and 13–3a.)

TB MED 576  
Occupational and Environmental Health: Sanitary Control and Surveillance of Water Supplies at Fixed Installations. (Cited in paras 12–2, 12–4, 13–3, and 13–4a.)

TB MED 577  
Occupational and Environmental Health: Sanitary Control and Surveillance of Field Water Supplies. (Cited in para 12–5b.)

TIM No. 14  
Protective Equipment for Pest Control Personnel. (Cited in para 10–14.) (This and the technical information memorandum listed below may be obtained from the Armed Forces Pest Management Board, Forest Glen Section, WRAMC, Washington, DC 20301–5001.)

TIM No. 21  
Pesticide Disposal Guide for Pest Control Shops. (Cited in para 10–13.)

TM 5–632/NAVFAC MO–310/AFM 9–16  
Military Entomology Operational Handbook. (Cited in paras 10–11 and 10–16.)

TM 5–666/AFR 91–26/NAVFAC MO–210  
Maintenance and Operation of Water Supply, Treatment, and Distribution Systems. (Cited in para 12–2f.)

TM 5–662  
Swimming Pool Operations and Maintenance. (Cited in paras 12–6a and 12–12c.)

Environmental Protection: Planning in the Noise Environment. (Cited in para 11–8a.)

TM 5–803–12  
Planning of Outdoor Recreation Areas. (Cited in para 12–12b.)

TM 5–810–1  
Mechanical Design: Heating, Ventilating, and Air Conditioning. (Cited in paras 9–7a(5) and 12–7d.)

TM 5–810–5/AFM 88–8  
Plumbing. (Cited in paras 12–7c and E–8d.)

TM 55–4470–400–12–1  
Transportability Guidance for Nuclear Reactor Irradiated Fuel Elements. (Cited in para 9–9b(13).)

UCMJ  

Unnumbered Publication  

Unnumbered Publication  
MIM’s Supplemental User’s Instructions. (Cited in para 5–7c.) (May be obtained from the Commander, USAEHA, ATTN: HSHB–MO–F, Aberdeen Proving Ground, MD 21010–5422.)

USAEHA TG No. 102  
Guide for the Conduct of Installation Pest Surveillance Programs. (Cited in paras 10–7b and c.) (This and the USAEHA technical guides listed below may be obtained from the Commander, USAEHA, ATTN: HSHB–CI–O, Aberdeen Proving Ground, MD 21010–5422.)

USAEHA TG No. 147  
Infectious Hazardous Waste Handling and Disposal. (Cited in para 11–7b(2).)

USAEHA TG No. 153  
Guidelines for Controlling Potential Health Hazards from Radiofrequency Radiation. (Cited in paras 9–4b(1), 9–5b, and 9–9a(2)(a) and (b) and b(12).)

USAEHA TG No. 167  
Hearing Evaluation Automated Registry System (HEARS) Audiometer Operation Manual. (Cited in para 5–16b(7)(e).)

10 CFR 20.103  
Standards for Protection Against Radiation: Exposure of individuals to concentrations of radioactive materials in air in restricted areas. (Cited in para 9–7a(1).)

29 CFR 1910  
Occupational Safety and Health Administration (OSHA) Standards. (Cited in paras 5–9a(1), 5–15c, 5–16b(9)(i),5–21b, and 8–4b.)

38 FR 21685  
Insecticides in Food Handling Establishments, Definitions and Policy Statements. (Cited in para 10–9a.)

Section II  
Related Publications

A related publication is merely a source of additional information. The user does not have to read it to understand this regulation.

ANSI A225.1–1984  
Manufactured Home Installation. (This and the ANSI publication listed below may be obtained from the American National Standards Institute Incorporated, 1430 Broadway, New York, NY 10018.)

ANSI Z16.4–1977  
Uniform Recordkeeping for Occupational Injuries and Illnesses.

AR 10–64/OPNAVINST 6700.2/AFR 160–29/MCO 5420.18A  
Joint Field Operating Agencies of the Office of The Surgeon General of the Army.

AR 11–2  
Internal Control Systems.

AR 40–2  
Army Medical Treatment Facilities: General Administration.

AR 40–3  
Medical, Dental, and Veterinary Care.

AR 40–10  
Health Hazard Assessment Program in Support of the Army Materiel Acquisition Decision Process.
AR 40–25/NAVMEDCOMINST 10110.1/AFR 160–95
Nutritional Allowances: Standards and Education.

AR 70–1
Systems Acquisition Policy and Procedures.

AR 190–24/MCO 1620.2/BUPERINST
Armed Forces Disciplinary Control Boards and Off-Installation Military Enforcement.

AR 190–38
Detention Cell Standards.

AR 190–47
The U.S. Army Correctional System.

AR 210–130
Laundry and Dry Cleaning Operations.

AR 600–85
Alcohol and Drug Abuse Prevention and Control Program.

AR 611–101
Personnel Selection and Classification, Commissioned Officer Classification System.

AR 635–100
Officer Personnel.

AR 690–700
Personnel Relations and Services (General).

AR 750–25
Army Test, Measurement, and Diagnostic Equipment (TMDE) Calibration and Repair Support Program.

C–3000/6300–IL
FSC Groups 30 through 63 (Items of Medical Materiel Only).

C–6700/9500–IL
Identification List: FSC Groups 67 thru 95 (Items of Medical Materiel Only).

DA Pam 600–17
A Commander’s, Supervisor’s, and Physician’s Guide to Alcohol and Alcoholism.

DA Poster 40–5
Lyme Disease Warning.

DOD Directive 4150.7
DOD Pest Management Program. (This and the DOD publications listed below may be obtained from the Commanding Officer, ATTN: Code 301, Naval Publications and Forms Center, 5801 Tabor Avenue, Philadelphia, PA 19120–5099.)

DOD Directive 6050.10
The Armed Forces Pest Management Board.

DOD Instruction 6050.5
Hazardous Material Information System.

DOD Instruction 6055.1
DOD Occupational Safety and Health Program.

DOD Instruction 6055.5
Industrial Hygiene and Occupational Health.

DOD Instruction 6055.12
DOD Hearing Conservation Program.

EPA 570/9–82–002
Manual for the Certification of Laboratories Analyzing Drinking Water. (This and the EPA publications listed below may be obtained from the National Technical Information Service, Port Royal Road, Springfield, VA 22161.)

EPA 600/4/79–019
Handbook for Analytical Quality Control in Water and Wastewater Laboratories.

EPA 600/8/78–017

FM 10–280
Mobile Field Laundry, Clothing Exchange, and Bath Operations.

FPM Chap 250
Personnel Management in Agencies.

FPM Chap 290
Personnel Information (General).

FPM Chap 293
Personnel Records and Files.

FPM Chap 294
Availability of Official Information.

FPM Chap 630
Absence and Leave.

FPM Chap 792
Federal Employees Health and Counseling Programs.

FPM Supp 792–2
Alcohol and Drug Abuse Programs.

FPM Chap 810
Injury Compensation.

FPM Chap 930
Programs for Specific Positions and Examinations.

GTA 8–5–45
Heat Injury Prevention and First Aid.

MIL STD 668
Sanitary Standards for Food Plants.

MIL STD 903
Sanitary Standards for Commissaries.

MIL STD 904

MIL STD 1486
In-Transit Fumigation of Freight Cars.

NFPA Std 56C
Safety Standards for Laboratories in Health Related Institutions. (This and the NFPA publications listed below may be obtained from the National Fire Prevention Association, Batterymarch Park, Quincy, MA 02269.)

NFPA Std 101

NFPA Std 491M
NFPA Std 501–A–1984
Mobile Home Installations, Sites, and Communities.

TB MED 288
Medical Problems of Man at High Terrestial Elevations.

TIM 24
Contingency Pest Management Pocket Guide. (May be obtained from the Armed Forces Pest Management Board, Forest Glen Section, WRAMC, Washington, DC 20301–5001.)

Children’s Play Areas.

TM 55–315

USAEHA TG No. 001
Appendixes E, G, and H for Medical Surveillance Guide, Guide for Job Related Examinations. (This and the USAEHA technical guides listed below may be obtained from the Commander, USAEHA, ATTN: HSHB–CI–O, Aberdeen Proving Ground, MD 21010–5422.)

USAEHA TG No. 028
Handling and Decontamination Guide for Elemental Mercury.

USAEHA TG No. 082

USAEHA TG No. 152

USAEHA TG No. 177

USAEHA TG No. 178

40 CFR 165
Regulations for the acceptance of certain pesticides and recommended procedures for the disposal and storage of pesticides and pesticides containers.

Section III
Prescribed Forms

DA Form 3075
Occupational Health Daily Log. (Prescribed in para 3–12.)

DA Form 3076

DA Form 3897–R
Tuberculosis Registry. (Prescribed in para 4–5h(14).)

DA Form 5402–R
Barber/Beauty Shop Inspection. (Prescribed in para D–8.)

DA Form 5931
Occupational Health Patient Form. (Prescribed in para 5–7.)

DA Form 5932
USAREUR Occupational Health Form. (Prescribed in para 5–7.)

DA Form 5933
Occupational Health Patient Form—Supplemental. (Prescribed in para 5–7.)

DA Form 5934
Korea Occupational Health Encounter Form. (Prescribed in para 5–7.)

DD Form 2215
Reference Audiogram. (Prescribed in para 5–21d(4).)

DD Form 2216
Hearing Conservation Data. (Prescribed in para 5–21d(5).)

DD Form 2493–1
Asbestos Exposure, Part I—Initial Medical Questionnaire. (Prescribed in para 5–28a.)

DD Form 2493–2
Asbestos Exposure, Part II—Periodic Medical Questionnaire. (Prescribed in para 5–28b.)

Section IV
Referenced Forms

DA Form 2417

DA Form 3761
Army Health Nursing Activities.

DA Label 80
U.S. Army Calibrated Instrument.

DD Form 689
Individual Sick Slip.

DD Form 1141
Record of Occupational Exposure to Ionizing Radiation.

DD Form 1532
Pest Management Report.

DD Form 2214
Noise Survey.

DOL Form CA 16
Authorization for Examination and/or Treatment.

DOL Form CA 17
Duty Status Report.

HEW Form CDC 73–2936A
Venereal Disease Epidemiologic Report.

HEW Form PHS 731
International Certificates of Vaccination.

OF 345
Physical Fitness Inquiry for Motor Vehicle Operators. (Replaces SF 47, which may be used.)

SF 78
United States Civil Service Commission Certificate of Medical Examination.

SF 93
Report of Medical History.

SF 177
Statement of Physical Ability for Light Duty Work.
A daily basis over a period of 1 to 2 weeks. ing in hot environments for gradually increasing periods of time on

tation to cold injury prevention by medical personnel.

chart in table 1, TB MED 81.

ative measures. All personnel should know how to use the wind chill

TB MED 81 describes the types of cold injuries and proper preven-

individual soldiers are essential to minimize cold injury casualties.

using RCS MED–16 under AR 40–400.

require hospitalization (such as heat exhaustion) will be reported

m e d i c a l  p e r s o n n e l . A  p o c k e t - s i z e d  g u i d e  t o  h e a t  i n j u r y  ( g r a p h i c

offered an annual orientation class on heat injury prevention by

prevention. The wet bulb globe temperature (WBGT) index and the

Botsball type, NSN 6665–01–103–8547, are available through supple-

portable WBGT Kit, NSN 6665–00–159–2218, and a WGT kit,

medical personnel should utilize at least one of these indexes during

degree of heat stress imposed by all environments. Commanders and

wet globe thermometer (WGT) are the best means of evaluating the

prevention. The wet bulb globe temperature (WBGT) index and the

TB MED 507 contains a comprehensive discussion of heat casualty

B–1. General

Climatic injuries of military importance include those disabilities to
troops caused by climatic or altitudinal factors.

B–2. Heat injuries

TB MED 507 contains a comprehensive discussion of heat casualty prevention. The wet bulb globe temperature (WBGT) index and the wet globe thermometer (WGT) are the best means of evaluating the degree of heat stress imposed by all environments. Commanders and medical personnel should utilize at least one of these indexes during all operations in heat, and especially for acclimatizing the troops. A portable WBGT Kit, NSN 6665–00–159–2218, and a WGT kit, Botsball type, NSN 6665–01–103–8547, are available through supply channels.

a. The following actions, if emphasized by the commander, will reduce the risk of heat injury:

(1) Acclimatization to heat. Acclimatization is acquired by working in hot environments for gradually increasing periods of time on a daily basis over a period of 1 to 2 weeks.

(2) Water intake. Adequate water intake is the single most important factor in avoidance of heat injury. An unlimited water drinking policy, particularly during times of increased physical stress, will be enforced.

(3) Salt. Undissolved salt tablets should never be used. There is sufficient salt in the American diet to preclude the use of supplemental salt solutions under normal circumstances. If salt supplements are required, salt solutions prepared as described in TB MED 507 should be given.

b. Each major unit should publish an annual directive on the prevention of heat injuries. Unit commanders and cadre should be offered an annual orientation class on heat injury prevention by medical personnel. A pocket-sized guide to heat injury (graphic training aid (GTA) 8–5–45) is available for individual use.

c. All heat injuries requiring hospital admission, or any significant clusters of heat injuries that occur in one unit that do not require hospitalization (such as heat exhaustion) will be reported using RCS MED–16 under AR 40–400.

B–3. Cold injuries

Careful prior planning and adequate training of commanders and individual soldiers are essential to minimize cold injury casualties. TB MED 81 describes the types of cold injuries and proper preventive measures. All personnel should know how to use the wind chill chart in Table 1, TB MED 81.

a. Unit commanders and cadre should be offered an annual orientation to cold injury prevention by medical personnel.

b. Cold injuries requiring hospital admission or any significant number of nonhospitalized cold injuries that occur in a unit will be reported using RCS MED–16 under AR 40–400.

B–4. High altitude

Acute mountain sickness may produce significant numbers of casualties for troops who are suddenly placed in a high altitude environment. Prevention is best achieved by thorough medical screening (AR 40–501) and acclimatization to altitude. A detailed discussion on high altitude is found in TB MED 288.

B–5. Requests for assistance

Commanders may request technical assistance on problems relating to heat, cold, and high altitude from the U.S. Army Research Institute of Environmental Medicine, Natick, MA 01760. Requests should be addressed through command channels to HQDA (SGPS–PSP), 5109 Leesburg Pike, Falls Church, VA 22041–3258.

Appendix C

Recommended Procedures for Cleanup of Pesticide Spills

C–1. General

These guidelines provide simple instructions on cleaning up small spills of pesticides (1 qt or less).

C–2. Personnel protection

a. If the pesticide gets into the eye or on the skin, immediately flush with water under low pressure. Further, the skin should be washed with soap and water. Remove contaminated clothing and blot up any pesticides on the clothing.

b. Once the pesticide is washed off and clothes changed, the persons should go to the nearest MTF. Exposed persons should know what pesticide they were exposed to and its strength or concentration.

c. Scoop up the contaminated absorbent and pour it into a plastic bag.

d. Mop up the spill area with warm water containing 1 cup of bleach (sodium hypochlorite) per gallon of water.

e. After mopping, place the mophead into the bag with the contaminated absorbent. Close and seal the bag and place it in a refuse container for pickup by an approved sanitation disposal service.

f. Cleanup personnel should wear coveralls and waterproof footwear (rubbers or galoshes). They should not eat, drink, or smoke during the cleanup and should wash thoroughly afterwards with soap and clean water.

g. Broken or damaged pesticide containers should be discarded in the same manner as the absorbent material.

Appendix D

Barber and Beauty Shop Sanitation

D–1. General

Skin disease agents may be transmitted either through direct contact or by fomites such as towels, combs, clippers, or razors. Skin diseases of concern include scalp ringworm (Tinea capitis), ringworm of the beard area of face and neck (Tinea barbae), and impetigo and staphylococcal infections.
D–2. Employee hygiene

a. Barbers and beauticians will not work when ill with communicable disease or other conditions which might be transferred to a patron.

b. The IMA will determine and state in a written policy if preemployment medical evaluations to ensure freedom from communicable disease, examinations before returning to work after illness, and special examinations are required. Examination authority, including incident hospitalization, is contained in AR 40–3.

c. Barbers and beauticians must keep their person and clothing clean when attending patrons. Smocks and/or uniforms will be changed daily.

d. Barbers and beauticians are prohibited from smoking, eating, and drinking in the work areas (such as in back bar areas, styling stations, and shampoo and drying areas). Smoking, eating, and drinking will be confined to designated employee break areas only. Customers are prohibited from smoking in barber and beauty shops.

D–3. Sanitary facilities

a. Barber and beauty shops will not be located in food service or sleeping areas.

b. Barber and beauty shops will be provided an adequate supply of hot and cold running water, proper plumbing fixtures, and adequate waste disposal. At fixed installations, a minimum of one lavatory will be provided for each two chairs. The lavatory will be located conveniently to both chairs served.

c. Shop interiors will be adequately lighted and ventilated.

d. Shops will be maintained in a sanitary condition at all times. Cut hair will be removed frequently from the floors.

e. Closed sanitary receptacles will be provided for waste materials and soiled linens.

D–4. Multiple service and disposable article sanitation(instruments, towels, and disposables)

a. Barber chair headrests will be covered with a clean sheet of paper or clean towel for each patron.

b. Freshly laundered towels or individual disposable sanitary neck strips will be used for each patron.

c. Reusable clean haircloths will be changed at least daily.

d. Use of common (natural bristle) brushes, neck dusters, shaving brushes, sponges, and powder puffs are prohibited. Excepted are synthetic hair brushes that are specifically designed to allow adequate cleaning and sanitizing between patrons. Use of automatic dispensers, brushless shaving cream, and clean towels in place of synthetic hair brushes that are specifically designed to allow adequate cleaning and sanitizing between patrons. Use of automatic dispensers, brushless shaving cream, and clean towels in place of disposable razors is recommended.

D–5. Sanitary practices

a. Without the written consent of a medical officer, patrons will not be served in barber or beauty shops when their face, neck, or scalp is inflamed, contains pus, or has erupted boils or pimples. Lice infested personnel will not be served and will be referred immediately for medical treatment.

b. Therapeutic practices such as treating blackheads, infected hairs, sores, or lesions are prohibited. Pulling of hairs from ears, nostrils, eyebrows, and mustaches is also prohibited.

c. Material used to stop blood flow will be in powder or liquid form, and should be applied with a freshly laundered towel or sterile absorbent cotton. Such material must be approved by a medical authority. The use of lump alum or styptic pencils is prohibited.

d. Caution should be exercised in the purchase and use of cosmetics, tonics, lotions, hair dyes, and bleaches. Some preparations have been implicated in skin and eye irritation and hair loss. All barber and beauty supplies must be approved by the USDA, FDA, or EPA for intended use.

D–6. Sanitization of instruments

a. Barbers and beauticians will clean all barbering instruments immediately after use on each patron. Scissors, combs, and tools will be thoroughly washed with soap and hot water to remove all film, oil, and debris, and then dried with a clean towel or clean disposable tissue.

b. Razors will be routinely disinfected between patrons to eliminate the possibility of transmission of infectious diseases such as hepatitis B. New disposable razors may be used on each patron if disinfection of reusable razors cannot be assured.

c. Removal of hair and debris from the exterior of clipper surfaces may be accomplished with a stiff bristle brush used only for this purpose.

d. If, in the course of a barbering process, it is suspected that a patron has a communicable disease or infection, the barbering instruments will be washed and disinfected immediately after use.

e. Instruments not intended to penetrate the skin, but which may become contaminated with blood (for example, razors), will be thoroughly cleaned and sanitized after use.

f. Disinfection will employ any liquid chemical disinfectant specifically formulated for barbering tools use and carrying a label registered by the USDA or EPA, or one approved by PVNTMED personnel. Germicides that are mycobacterial are preferred because mycobacteria are one of the most resistant groups of microorganisms. Disinfectants will be used according to label instructions. Other disinfection procedures, such as ultraviolet, will be used only with medical authority approval. Disinfection solutions will be prepared and changed frequently enough to ensure bactericidal effectiveness when used or at least once daily.

g. Containers for instrument disinfection will be provided with covers and be of sufficient size to accommodate all instruments.

h. At the close of each day’s operation, all barbering tools used will be washed and disinfected.

i. All barbering instruments disinfected in a chemical solution will be rinsed in running water to remove chemicals before patron use.

D–7. Posting of regulation

A copy of this appendix will be maintained (preferably in a folder on a magazine rack) for customer inspection in each barber and beauty shop.

D–8. Inspection form

a. DA Form 5402–R (Barber/Beauty Shop Inspection) will be locally reproduced on 8 1/2 by 11-inch paper. DA Form 5402–R is located at the back of this regulation.

b. Requirements on this form are directly related to requirements in this appendix. Use of this form is strongly recommended for all PVNTMED services.

D–9. Field barber’s kit

All military personnel using the field barber’s kit (national stock number 3590–00–058–1837) at organizational and unit levels will sanitize all barbering instruments before and after each use per paragraph D–6.

Appendix E

Mobile Home Parks Sanitation

E–1. General

Mobile home parks include locations intended for permanent or semipermanent places of residence. They do not include locations of temporary residence intended for recreational vehicles, travel trailers, and similar vehicles.

E–2. Location

Mobile home parks will be located in level, well drained areas and should not be located adjacent to swamps, marshes, breeding places for insects and rodents, or heavy industrial zones. The mobile home park should have good natural drainage, or a storm drainage system
must be provided. Storm drainage must not endanger any water supply. All-weather roads, both to and within the park, will be provided.

**E–3. Individual parking areas**

Each area will be at least 45 by 70 feet and surfaced to provide a level, well-drained space under and adjacent to the mobile home. In mobile home parks that allow parking of double wide or extended length mobile homes, minimum individual parking areas for these trailers will be at least 25 feet wider and 20 feet longer than the trailer.

**E–4. Mobile home**

The mobile home will be of substantial construction and designed and constructed according to standards of commercial-type trailers. At least 35 square feet of floor space per occupant will be provided. Lean-tos, sheds, or additional rooms will not be attached to the mobile homes. Open porches, awnings, and original equipment expandable rooms are authorized, provided a minimum clear area of 10 feet between the mobile home and the individual parking area line is maintained. If locally authorized, centralized or individual storage sheds may be erected, provided they are equipped with suitable foundations and floorings and are not used for human habitation.

**E–5. Water supply**

Potable water will be provided at each mobile home space by means of suitable sanitary connections. Plumbing and sewage will be designed and installed under the current National Standard Plumbing Code.

**E–6. Liquid waste and wash water disposal**

A vertical drain pipe equipped with a suitable trap and connected to a sanitary sewer will be provided at each mobile home space. The connection between the drainage system of the mobile home and the vertical drain will be made to exclude insects and rodents, prevent leakage and escaping odors, and otherwise prevent health hazards.

**E–7. Human waste disposal**

The mobile home water closet connection will only be made by facility engineer personnel and then only when—

- a. Mobile home plumbing fixtures and the system are approved by the facilities engineer and the medical authority.
- b. The mobile home park sewer system is designed, installed, and operated under Army standards. Liquid wastes will drain into an approved sewer system and/or treatment and disposal facility. Requests for exception will be submitted to HQDA (SGPS–PSP), 5109 Leesburg Pike, Falls Church, VA 22041–3258.

**E–8. Service buildings**

Each mobile home park will have at least one service building to provide necessary sanitation and laundry facilities.

- a. Heating facilities will be capable of maintaining a temperature of 65 degrees Fahrenheit in cold weather.
- b. Adequate lighting will be provided inside and outside buildings.
- c. Service buildings will be conveniently located within 100 yards of the most remote mobile home space.
- d. Every mobile home park will provide adequate toilet and laundry facilities as indicated in TM 5–810–5/AFM 88–8. These fixtures are necessary to provide adequate facilities when mobile homes are repaired, connected, disconnected, and/or used for other emergencies, even though the mobile home park may accommodate only independent coaches.

**E–9. Area sanitation**

Roads, car parks, sidewalks, and other areas will be provided with surfacing to control dust and mire. Adequate drainage will be provided to prevent accumulations of surface water.

**E–10. Illumination and fire protection**

Adequate area illumination and fire protection will include a suitable electrical outlet at each mobile home space. Area illumination should be arranged to avoid mobile home occupant annoyance.

**E–11. Protection of utility connections**

Mobile home utility terminals will be adequately secured. Terminals will be located to assure protection from tampering, breakage, or contamination.

**E–12. Design criteria**

Design criteria for development and evaluation of mobile home parks are provided in ANSI A225.1–1984 and NFPA Standard 501–A–1984. These publications, in addition to the specific requirements of this regulation, should be used in developing local mobile home park sanitation programs as required.
Glossary

Section I
Abbreviations

AFAP
Army family advocacy program

AFPMB
Armed Forces Pest Management Board

AHS
Academy of Health Sciences, U.S. Army

AMEDD
Army Medical Department

AMH
Accreditation Manual for Hospitals (JCAHO)

ANSI
American National Standards Institute

AOC
area of concentration (formerly SSI: specialty skill identifier)

ARD
acute respiratory disease

BCG
Bacille Calmette-Guerin

CDS
child development services

CFR
Code of Federal Regulations

CHEMTREC
Chemical Transportation Emergency Center

CHN
community health nurse

CHR
command health report

CINCUSARPAC
Commander in Chief, U.S. Army, Pacific

CONUS
continental United States

CTA
Common Table of Allowances

DA
Department of the Army

DENTAC
dental activity

DHS
director, health services

DLA
Defense Logistics Agency

DOD
Department of Defense

DOL
Department of Labor

DPCCM
department of primary care and community medicine

EMSL–CI
Environmental Monitoring and Support Laboratory, Cincinnati, Ohio

EPA
Environmental Protection Agency

EPICON
epidemiology consultant (assistance) (service)

FACMT
family advocacy case management team

FC
field circular

FDA
Food and Drug Administration

FM
field manual

fpm
feet per minute

FPM
Federal Personnel Manual

GTA
graphic training aid

HHA
health hazard assessment

HHIM
Health Hazard Information Module

HIV
human immunodeficiency virus

HQDA
Headquarters, Department of the Army

HSC
Health Services Command (U.S. Army)

IMA
installation medical authority

INH
isoniazid

IPM
integrated pest management

JCAHO
Joint Commission on Accreditation of Healthcare Organizations

kg
kilogram(s)

m
meter

MACOM
major Army command

MADP
materiel acquisition decision process

MEDCAC
U.S. Army medical department activity

MEDDAC
U.S. Army medical center

MIL STD
military standard

mg
milligram(s)

MIM
Medical Information Module

min
minute

mm
millimeter(s)

m/min
meter(s) per minute

MTF
medical treatment facility

NBSH
National Bureau of Standards handbook

NFPA
National Fire Prevention Association

NIOSH
National Institute for Occupational Safety and Health

NPDR
National Primary Drinking Water Regulation

NRC
Nuclear Regulatory Commission

OCONUS
outside continental United States

OH
occupational health

OHE
occupational health education

OHIMS
Occupational Health Management Information System

OPM
Office of Personnel Management

**Note**: Section III of this glossary contains copyright material. For notice, see cover.
Definitions

**OSHA**
O c c u p a t i o n a l  S a f e t y  a n d  H e a l t h  A d m i n i s t r a t i o n

**OTSG**
Office of the Surgeon General

**PCS**
permanent change of station

**pH**
hydrogen ion concentration

**PMM**
preventive medicine measures

**POC**
point of contact

**PPE**
personal protective equipment

**ppm**
parts per million

**PVNTMED**
preventive medicine

**QA**
quality assurance

**qt**
quart

**RCS**
Requirement Control Symbol

**rem**
roentgen equivalent man

**RPO**
radiation protection officer

**SOH**
safety and occupational health

**SOP**
standing operating procedure

**STD**
sexually-transmitted disease

**TDA**
table(s) of distribution and allowances

**TG**
technical guide

**TIM**
technical information memorandum

**TM**
technical manual

**TMDE**
test, measurement, and diagnostic equipment

**TOE**
table(s) of organization and equipment

**TSG**
The Surgeon General

**TST**
tuberculin skin test

**UCMJ**
Uniform Code of Military Justice

**USAEHA**
U.S. Army Environmental Hygiene Agency

**USAPACEHEA**
U.S. Army Pacific, Environmental Health Engineering Agency

**USAHEUR**
U.S. Army, Europe

**USDA**
U.S. Department of Agriculture

**USPHS**
U.S. Public Health Service

**VDT**
video display terminal

**WBGT**
wet bulb globe temperature

**WGT**
wet globe thermometer

**Section II**
Terms

**Disposal**
The discharge, deposit, injection, dumping, spilling, leaking, or placing of any solid waste or hazardous waste into or on any land or water so that such solid waste or hazardous waste or any constituent thereof may enter the environment or be emitted into the air or discharged into any waters, including ground waters.

**General waste**
All healthcare facility waste not classified as infectious, pathological, or hazardous. Examples may include refuse generated in general patient units, emergency rooms, dental areas, surgical suites, administrative areas, and supply areas.

**Hazardous waste**
Solid waste, or combination of solid waste (except those excluded in 40 CFR 261.4(b)), that because of its quantity, concentration, or physical, chemical, or infectious characteristics may—

a. Cause, or significantly contribute to an increase in mortality or an increase in serious, irreversible or incapacitating, reversible illness.

b. Pose a substantial present or potential hazard to human health or the environment when improperly treated, stored, transported, or disposed of, or otherwise managed.

**Healthcare facility**
A structure or location where patient care is provided by AMEDD personnel.

**Incineration**
A method of thermal treatment of general infectious or pathological waste.

**Industrial hygiene**
The OH science consisting of the recognition, evaluation, and control of workplace health hazards.

**Infectious waste**
Infectious waste is any waste with pathogens of sufficient virulence and quality capable of causing an infectious disease in an exposed susceptible host. These wastes are generally from patients in strict or respiratory isolation, or with wound and skin precautions; wastes from the microbiology laboratory; surgical waste at the discretion of the infection control committee. State/local definitions may be more restrictive and would then prevail.

**Installation medical authority**
The unit surgeon, command chief surgeon, MEDDAC and/or MEDCEN commanders, and the DHS, or his or her representative responsible for provision of medical support at the unit, command, or installation concerned.

**Occupational medicine staff**
Personnel who may include an OH physician, OH nurse, industrial hygienist, industrial hygiene/OH technician, RPO, audiologist, and optometrist.

**Pathological waste**
Any wastes that include anatomical parts of humans and animals, excluding human corpses and animal carcasses. State/local definitions may be more restrictive and would then prevail.

**Permitted sanitary landfill**
A landfill that has State or Federal approval to operate and is operated in a manner that protects health and the environment. Waste is compacted and covered with earth daily, scavenging is strictly prohibited, and it is not an attractant to vermin.

**Refuse**
All solid waste that is not classified as infectious, pathological, or hazardous waste. Often referred to as trash and garbage.

**Respiratory isolation**
Isolation that prevents transmission of organisms by means of direct contact or droplets that are coughed, sneezed, or breathed into the environment.

**Retort**
A large steam autoclave, constructed usually
of material other than stainless steel, used to treat infectious or pathological waste.

**Solid waste**

Any garbage, refuse, or sludge from a waste treatment plant, water supply treatment plant, or air pollution control facility. It includes other discarded material such as solid, liquid, semisolid, or contained gaseous material resulting from industrial, commercial, mining, and agricultural operations, and from community activities. Infectious, pathological, and hazardous wastes are special categories of solid waste. Each of these categories requires specific management in collection, handling, and disposal. The following are not solid wastes—

a. Solid or dissolved material in domestic sewage.

b. Solid or dissolved materials in irrigation return flows.

c. Industrial discharges which are point sources subject to permits under section 402 of PL 92–500 (The Federal Water Pollution Control Act of 1972) as amended.


**Strict isolation**

Isolation that prevents the transmission of all highly communicable diseases that are spread by both contact and airborne routes of transmission.

**Treatment of hazardous waste**

Any method, technique, or process, including neutralization, designed to change the physical, chemical, or biological character or composition of any hazardous waste that neutralizes such waste or renders such waste nonhazardous, safer for transport, amenable for recovery, amenable for storage, or reduced in volume. Such term includes any activity or processing designed to change the physical form or chemical composition of hazardous waste to render it nonhazardous.

**Workplaces**

a. Nonmilitary-unique workplaces and operations. DOD military and civilian workplaces and operations generally comparable to those of business and industry in the private sector. Examples include facilities involved and work performed in the repair and overhaul of weapons, vessels, aircraft, or vehicles (except for equipment trials); construction; supply services; civil engineer or public works; medical services; and office work.

b. Military-unique equipment, systems, operations, or workplaces.

(1) Equipment and systems that are unique to the National defense mission, including the operation, testing, and maintenance procedures dictated by the design configuration. Examples include military weapons, aircraft, ships, submarines, missiles and missile sites, early warning systems and sites, military space systems, ordnance, tanks, and tactical vehicles.

(2) Operations or workplaces that are uniquely military, such as field maneuvers; combat training; naval operations; military flight and missile operations; associated research, test, and development activities; and actions required under emergency conditions. Wound and skin precautions Precautions that prevent acquisition of infection by personnel and patients from direct contact with wounds and heavily contaminated articles.

**Section III**

Definitions Extracted from ANSI Standard Z16.41977

(This Appendix is not an official part of American National Standard for Uniform Recordkeeping for Occupational Injuries and Illnesses, Z16.41977, but is included within that publication for information purposes only.)

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**Guiding Interpretations and Examples**

(The subdivisions of the appendix are numbered to correspond with the section numbers in the standard. Since it is not necessary to have appendix matter for every section in the standard, there are gaps in the numbering in the appendix.)

**A2. Definitions**

A2.2 Exposure or Employee-Hours. Determination of employee-hours of exposure: Employee-hours of exposure for calculating incidence rates are to be actual hours worked. When actual hours are not available, estimated hours may be used. Employee-hours should be calculated as given under A2.2.1 and A2.2.2.

A2.2.1 Actual Exposure Hours. Employee-hours of exposure for nonexempt employees are to be taken from payroll or time-clock records and include only actual straight-time hours worked and actual overtime hours worked.

A2.2.2 Estimated Exposure Hours. When actual employee-hours of exposure are not available, estimated hours may be used. Such estimated hours should be obtained by multiplying the total employee-days worked for the period by the average number of hours worked per day. If the hours worked per day vary among departments, a separate estimate should be made for each department and these estimates added to obtain the total hours. Estimates of overtime hours should be included.

Note. If employee-hours are estimated, indicate the basis on which estimates are made.

A2.2.2.1 Employees Living on Company Property. In calculating hours of exposure for employees who live on company property, only those hours during which employees were actually on duty are to be counted.

A2.2.2.2 Employees with Undefined Hours of Work. For traveling personnel, executives, and others whose working hours are not defined, an average of 8 hours per day is to be assumed in computing exposure hours.

A2.2.2.3 Standby Employees. For standby employees, including seamen aboard vessels, who are restricted to the confines of the employer's premises, all standby hours must be counted.

A2.6 Medical Treatment. Medical treatment versus first aid: The important point to be stressed is that the decision as to whether a case involves medical treatment should be made on the basis of whether the case normally would require medical treatment. The decision cannot be made on the basis of who treats the case. First aid can be administered by a physician and medical treatment by someone other than a physician.

It is not possible to list all types of medical procedures and treatments and on that basis alone determine if first aid or medical treatment was involved. For example, whirlpool treatments, heat treatments, application of hot or cold compresses, or elastic bandages are not in and of themselves either first aid or medical treatment.

What follows is a discussion of diagnostic procedures and preventive procedures and treatments, both of which are not in and of themselves medical treatment. Next is a discussion of treatments that are almost always medical treatment, and comments on medical treatment and first aid for certain types of injuries.

**A2.6.1. Diagnostic Procedures**

A2.6.1.1 Hospitalization for observation, where no medical treatment is rendered other than first aid, is not considered medical treatment.

A2.6.1.2 Visits to a physician or nurse for observation only or for routine change of dressing are not considered medical treatment.

A2.6.1.3 X-ray examination for fractures is considered diagnostic procedure and as such is not considered medical treatment or first aid. Where the x-ray is negative, the case is not recordable unless the injury required other medical treatment or met one of the other criteria for recordability.

A2.6.1.4 Physical examination yielding few or no findings and not substantiating subjective complaints in questionable cases is not considered medical treatment.

A2.6.1.5 Reactions to or effects of diagnostic procedures that are necessitated by a work-related injury or illness and which meet the criteria for recordability should be recorded.
A2.6.2. Preventive Procedures and Treatments
A2.6.2.1 Tetanus shots, either initial shots or boosters, are considered preventive in nature and are not in and of themselves considered medical treatment. However, treatment of a reaction to a tetanus shot administered because of an injury would be considered medical treatment and would make the case recordable.

A2.6.2.2 Any use of prescription medication normally constitutes medical treatment. However, it should be considered first aid when a single dose or application of a prescription medication is given on the first visit merely for relief of pain or as preventive treatment for a minor injury. This situation can occur at facilities having dispensaries stocked with prescription medications frequently used for preventive treatment and relief of pain and attended by a physician or registered professional personnel operating under the standing orders of a physician. The administration of nonprescription medication in similar circumstances would be considered first aid.

A2.6.2.3 The application of ointments and salves to prevent the drying or cracking of skin at the site of a minor injury can be considered first aid.

A2.6.2.4 The application of antiseptics to minor injuries which do not themselves require medical treatment can be considered first aid. Changing the bandage or dressing on an injury which did not require medical treatment, because the bandage or dressing has become dirty, is considered first aid.

A2.6.2.5 Reaction to preventive medication (not administered because of an occupational injury or illness) administered in-plant (such as flu shots) would not constitute a recordable case.

A2.6.2.6 In-plant treatment of off-the-job injuries and illnesses is not recordable.

A2.6.3 Treatments That Are Almost Always Medical Treatment: These treatments are as follows:
(1) Suturing of any wound
(2) Treatment of fractures
(3) Application of a cast or other professional means of immobilizing an injured part of the body
(4) Treatment of infection arising out of an injury
(5) Treatment of a bruise by the drainage of blood
(6) Surgical debridement, that is, the removal of dead or damaged tissue
(7) Treatment of abrasions that occur to greater than full skin depth
(8) Treatment of second- and third-degree burns

Note: Administration of prescription medicines is usually considered medical treatment (see A2.6.2.2).

A2.6.4. Medical Treatment and First Aid for Certain Types of Injuries

A2.6.4.1. Cuts and Lacerations
A2.6.4.1.1 First Aid. Treatment is limited to cleaning of the wound, soaking, application of antiseptic or nonprescription medication, and bandaging on the first visit. Follow-up visits are limited to observation, including changing of the dressing and bandage. Additional cleaning and application of antiseptic are permissible as first aid where required by work duties that are likely to soil the bandage. Application of butterfly closures for cosmetic purposes only can be considered first aid.

A2.6.4.1.2 Medical Treatment. The injury requires butterfly closures (for noncosmetic purposes), sutures (stitches), surgical debridement (cutting away dead tissue), treatment of infection, or other professional treatment.

A2.6.4.2. Abrasions
A2.6.4.2.1 First Aid. This is the same as for cuts and lacerations except that ointments can be added on follow-up visits to prevent drying and cracking of skin.

A2.6.4.2.2 Medical Treatment. The injury requires careful examination for removal of embedded foreign material, multiple soakings, whirlpool treatment, treatment of infection, or other professional treatment. This is any case involving more than a minor, spot-type injury. Treatment of abrasions occurring to greater than full skin depth is considered medical treatment.

A2.6.4.3. Bruises
A2.6.4.3.1 First Aid. Treatment is limited to a single soaking or application of cold compresses on a minor bruise. Follow-up visits are limited only to observation.

A2.6.4.3.2 Medical Treatment. The injury requires multiple soakings, draining of collected blood, or other extended care beyond observation.

A2.6.4.4. Splinters and Puncture Wounds
A2.6.4.4.1 First Aid. Treatment is limited to cleaning of the wound, removal of a foreign object(s) by tweezers or other simple techniques, application of antiseptics and nonprescription medications, and bandaging on the first visit. Follow-up visits are limited to observation, including changing of the bandage. Additional cleaning and application of antiseptic are permissible as first aid where required by work duties that are likely to soil the bandage.

A2.6.4.4.2 Medical Treatment. The injury requires removal of a foreign object(s) by a physician due to the depth of embedment, size or shape of the object(s), or location of the wound. This is also injuries requiring treatment for infection, treatment of a reaction to a tetanus booster, or other professional treatment.

A2.6.4.5. Burns, Thermal and Chemical (Resulting in Destruction of Tissue by Direct Contact)
A2.6.4.5.1 First Aid. Treatment is limited to cleaning or flushing of the surface; soaking; application of cold compresses, antiseptics, or nonprescription medications; and bandaging on the first visit. Follow-up visits are restricted to observation, changing of bandages, or additional cleaning. Most first-degree burns are amenable to first-aid treatment.

A2.6.4.5.2 Medical Treatment. The injury requires a series of treatments including soaks, use of whirlpools, and surgical debridement (cutting away dead tissue). Most second- and third-degree burns require medical treatment.

A2.6.4.6. Sprains and Strains
A2.6.4.6.1 First Aid. Treatment is limited to soaking, application of cold compresses, and use of an elastic bandage on the first visit. Follow-up visits are for observation, possibly including reapplying a bandage.

A2.6.4.6.2 Medical Treatment. The injury requires a series of hot and cold soaks, use of whirlpools, diathermy treatment, or other professional treatment.

A2.6.4.7. Eye Injuries
A2.6.4.7.1 First Aid. Treatment is limited to irrigation, removal of foreign material not embedded in the eye, and application of nonprescription medications. A precautionary visit (special examination) to a doctor is still considered first aid if treatment is limited to the aforementioned items. Follow-up visits are for observation only.

A2.6.4.7.2 Medical Treatment. This is cases involving removal of embedded foreign objects, use of prescription medications, or other professional treatment.

A2.6.4.8. Inhalation of Toxic or Corrosive Gases
A2.6.4.8.1 First Aid. Treatment is limited to removal of the employee to fresh air or the one-time administration of oxygen for several minutes.

A2.6.4.8.2 Medical Treatment. This is any professional treatment beyond the aforementioned. It includes all cases involving loss of consciousness.

A2.7 Work-Related Cases. The broad concept is that any injury or illness “occurring in and attributable to the work environment” is “work-related.” Work environment is comprised of the physical location, equipment, materials processed or used, and the kinds of operations performed by an employee in the performance of his work, whether on or off the employer’s premises. There are no stated exclusions of place or circumstance. Therefore, injuries or illnesses occurring in such places as the employee parking lot, lunchroom, or restroom, or during rest or lunch period on the employer’s premises, can be work-related. The final determination of whether any case is work-related must be made by the employer. Responsibility or fault does not enter into the decision of whether a
case is work-related. In doubtful situations, a case should be recorded.

A2.8. Recordable Cases
Recordable work-related cases are those that involve any of the following:

(1) Deaths, regardless of the time between occupational injury or illness and death.

(2) All occupational illnesses, including, but not limited to, the following categories and examples:

(a) Occupational skin diseases or disorders
Examples: contact dermatitis, eczema, or rash caused by primary irritants and sensitizers, or poisonous plants; oil acne; chrome ulcers; chemical burns or inflammations. (Direct contact causing tissue damage only, resulting from a thermal or chemical burn, is classified as an injury, not an illness case.)

(b) Dust diseases of the lungs (pneumoconioses)
Examples: silicosis, asbestosis, coal worker’s pneumoconiosis, byssinosis, and other pneumoconioses.

(c) Respiratory conditions due to toxic agents
Examples: pneumonitis, pharyngitis, rhinitis or acute congestion due to chemicals, dusts, gases or fumes; farmer’s lung.

(d) Poisoning (systemic effects of toxic materials)
Examples: poisoning by lead, mercury, cadmium, arsenic, or other metals; poisoning by carbon monoxide, hydrogen sulfide, or other gases; poisoning by benzol, carbon tetrachloride, or other organic solvents; poisoning by insecticide sprays such as parathion, lead arsenate; poisoning by other chemicals such as formaldehyde, plastics, and resins.

(e) Disorders due to physical agents (other than toxic materials)
Examples: heatstroke, sunstroke, heat exhaustion, and other effects of environmental heat; freezing; frostbite and effects of exposure to low temperatures; caisson disease; effects of ionizing radiation (x-rays, radium); effects of nonionizing radiation (welding flash, ultraviolet rays, microwaves, sunburn).

(f) Disorders associated with repeated trauma
Examples: noise-induced hearing loss; synovitis, tenosynovitis, and bursitis; Raynaud’s phenomenon; and other conditions due to repeated motion, vibration, or pressure.

(g) All other occupational illnesses
Examples: anthrax, brucellosis, infectious hepatitis, malignant and benign tumors, food poisoning, histoplasmosis, coccidioidomycosis.

(3) Injuries resulting in any of the following:

(a) Lost workday—either days away from work or days of restricted work activity
(b) Medical treatment other than first aid
(c) Loss of consciousness
(d) Restriction of work or motion
(e) Temporary or permanent transfer
(f) Termination of injured or ill employee
Loss of consciousness of the employee for any period of time is self-explanatory. Restriction of motion is not defined specifically. Each case must be judged individually to determine if there is more than a trivial amount of restricted motion, such as would occur when a small adhesive bandage was placed on the second joint of the finger. It should be noted here that damage to prostheses (such as false teeth) is not in and of itself grounds for recordability unless accompanied by other damage to the body that meets the recordability criteria.

A2.10.1 Lost Workday Cases with Days Away from Work.
These are cases that result in 1 or more days away from work. Days away from work are those workdays (consecutive or not) on which the employee would have worked but could not because of occupational injury or illness. The number of lost workdays should not include the day of injury or onset of illness or any days on which the employee would not have worked even though able to work. For example, if an employee who is scheduled to work Monday through Friday has a recordable case on Friday and returns to work on Monday, the case does not involve any days away from work even if the employee was unable to work on Friday. Saturday, or Sunday. If this same employee had been scheduled to work on Saturday, even if that Saturday constituted overtime, the Saturday would be counted as days away from work, and the case would be classified as a lost workday case with days away from work.

For employees not having a regularly scheduled shift, for example, certain truck drivers, construction workers, farm labor, casual labor, part-time employees, etc., it may be necessary to estimate the number of lost workdays. Estimates of the number of days that the employee would have worked should take into account the prior work history of the employee and days worked by employees, not ill or injured, working in the same department or occupation as the ill or injured employee. In some cases an injured or ill employee will miss one or more scheduled days or shifts besides the day of injury or onset of illness, but it will be uncertain whether the employee was truly unable to work on the days missed. Such cases may arise when a doctor judges that the employee is able to work but the employee decides that he is not. In such cases, the employer should not rely solely on the doctor’s opinion. He should make the final judgement himself based on all the evidence at his disposal. Again, the rule should be “when in doubt, record the case.”

A2.10.2 Lost Workday Cases with Days of Restricted Work Activity Only.
These are cases that result in 1 or more days of restricted work activity but do not result in any days away from work.

Days of restricted work activity include those days (consecutive or not, but excluding the day of injury or onset of illness) on which
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<td>DRUG REGIMEN</td>
<td></td>
</tr>
<tr>
<td>DATE STARTED</td>
<td>TYPE</td>
</tr>
</tbody>
</table>


DA FORM 3897-R, AUG 90 Previous editions obsolete. **TUBERCULOSIS REGISTRY**

For use of this form, see AR 40-5; the proponent agency is the OTSG
REPORTED TO STATE/LOCAL HEALTH DEPARTMENT (If yes, list name of State or Local Health Department and date reported.)

☐ YES ☐ NO

X-RAY FINDINGS

<table>
<thead>
<tr>
<th>DATE</th>
<th>TEST RESULTS</th>
<th>DATE</th>
<th>NOTES</th>
</tr>
</thead>
</table>

RETURN VISIT ACTIONS (Use pencil).

Reverse of DA Form 3897-R, AUG 90
## BARBER/BEAUTY SHOP INSPECTION

For use of this form, see AR 40-5, the proponent is TSG

### INSTALLATION

<table>
<thead>
<tr>
<th>BUILDING NO.</th>
<th>FACILITY DESIGNATION</th>
</tr>
</thead>
</table>

### PERSON IN CHARGE OF FACILITY

<table>
<thead>
<tr>
<th>TYPE FACILITY</th>
<th>COPY REPORT FURNISHED TO</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. AAFES Barber Shop</td>
<td>1. Satisfactory</td>
</tr>
<tr>
<td>2. Troop Barber Shop</td>
<td>2. Unsatisfactory</td>
</tr>
<tr>
<td>3. Club Barber Shop</td>
<td>3. Marginal</td>
</tr>
<tr>
<td>6. Other (specify)</td>
<td>4. Other (specify)</td>
</tr>
</tbody>
</table>

### PURPOSE

1. Regular
2. Courtesy
3. Pre-opening
4. Other (specify)

### COMMAND INSTALLATION FACILITY INSPECTOR INSPECTION DATE (DAY) YEAR NO. DAY RESERVED

| 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 | 16 | 17 | 18 | 19 | 20 | 21 | 22 | 23 | 24 | 25 | 26 | 27 | 28 | 29 | 30 | 31 | 32 | 33 | 34 |

### Description

(Use reverse for remarks; identify each remark by item number.)

| Reference Paragraph ARI 40-5 | Deficiency Yes | No |

### Employee Hygiene

1. Employees do not work when ill with communicable disease (e.g., boils, skin infections, etc.)

2. Current preemployment or periodic medical examination certificates (only when required by medical authority)

3. Employees do not return to work after illness unless cleared by the medical authority

4. Clean uniform worn

5. Employees smoke, eat or drink only in designated break areas

### Sanitary Facilities

6. Not located in food service or sleeping areas

7. Adequate hot and cold running water, adequate fixtures and waste disposal, no cross connections

8. Shop area kept clean, adequately lighted and ventilated. Outside area policed

9. Adequate closed waste receptacles provided

### Instruments, Towels, and Disposables

10. Headrest covered with clean paper or towel for each patron

11. Only individual freshly laundered or disposable neck strips used

12. Reusable haircloth kept clean and changed at least daily

13. No common brushes, neck dusters, shaving brushes or other similar multiuse brushes used. (Exception allowed for synthetic bristle brushes which are designed to be cleaned between patrons and sanitized as required.)

### Sanitary Practice

14. Patrons with boils, pimples or other inflammations referred to medical authority prior to services

15. Persons with known or suspected skin infections not served

16. Prohibited practices not conducted (e.g. treating blackheads, removing hairs from ears, nose, etc.)

17. Only approved materials used for stopping blood

18. Materials applied with freshly laundered or disposable cloths. Cloths disposed of properly

19. Only USDA, FDA, or EPA approved barber and beauty supplies used and only for intended use

### Sanitation of Instruments

20. Instruments scrupulously cleaned between patrons

21. Hair removed from clippers between patrons

22. Instruments disinfected as required

23. Only USOA or EPA disinfectants used. Disinfectants used in accordance with label instructions

24. Fresh solution prepared at least daily

25. All instruments rinsed with potable water after disinfecting

26. Copy of Appendix E, ARI 40-5, posted

### Critical deficiencies requiring immediate correction

- **Sanitary Practice**
- **Sanitation of Instruments**

### Inspected By

(Signature)

Date/Time

Copy Received By

(Signature and date)

DA FORM 5402-R, MAR 85