

CE Activity Announcement

Sterile Product Preparation Certificate

ACPE Activity Number: 0204-0000-19-725-H03-P & T thru to 0204-0000-19-738-H07-P & T

Release Date: April 10, 2019

Expiration Date: April 10, 2022

Activity Type: Application-based

CE Credits: 29.00 hours (*no partial credit*)

Accreditation for Pharmacists and Pharmacy Technicians



The American Society of Health-System Pharmacists is accredited by the Accreditation Council for Pharmacy Education as a provider of continuing pharmacy education.

Target Audience

This program is intended for pharmacy technicians and pharmacists engaging in safe and compliant sterile product preparation.

Activity Overview

The program provides 29 hours of ACPE continuing education for pharmacists and technicians, incorporating recorded presentations, readings, video demonstrations, and exercises in curricular modules. The program covers sterile product preparation practice standards and regulations, pharmacy calculations, facilities and equipment, cleanroom personnel behaviors and expectations, sterile compounding components and procedures, stability and sterility, nonsterile to sterile compounding, hazardous drugs, and all aspects of handling from material receipt to final check or disposal. After completing all of the modules, participants should be proficient in the fundamental concepts required to ensure safe and compliant sterile product preparation.

Learning Objectives and Schedule of Activities

Activity CE Information	Title, Description, and Learning Objectives
<p>ACPE #: 0204-0000-19-725-H03-P 0204-0000-19-725-H03-T</p> <p>CE Hours: 1.25 Activity Type: Knowledge</p>	<p>Title: Introduction to Sterile Product Preparation</p> <p>This activity discusses the guidelines, best practices, standards, and regulations that influence and contribute to safe sterile product preparation practices.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> • Identify factors contributing to the development of sterile compounding guidelines, best practices, standards, and regulations in the United States. • Describe lessons learned from major patient safety events involving sterile compounded preparations. • List the standards of practice that apply to sterile compounding in the United States. • Identify best practices to ensure sterile compounding safety. • Explain the role of the compounder in assuring the safety of compounded sterile preparations.
<p>ACPE #: 0204-0000-19-726-H07-P</p>	<p>Title: Pharmacy Calculations</p>

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<p>0204-0000-19-726-H07-T</p> <p>CE Hours: 1.75 Activity Type: Application</p>	<p>This activity covers calculations routinely used by pharmacy personnel when compounding sterile preparations.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> • Calculate doses in both weight and volume using ratios, proportions, and concentrations. • Practice conversions commonly used in sterile compounding calculations. • Apply the alligation method to calculate parts of two solutions with different concentrations to compound a solution with a different desired concentration. • Calculate infusion times in mL/hr and mL/min. • Select quantity of dosage units required to supply an order for a specified number of days or weeks.
<p>ACPE #: 0204-0000-19-727-H07-P 0204-0000-19-727-H07-T</p> <p>CE Hours: 1.5 Activity Type: Application</p>	<p>Title: Overview of Compounding Facilities and Engineering Controls</p> <p>This activity explains facility requirements and how primary and secondary engineering controls support sterile product preparation.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> • Describe sterile compounding facility requirements. • Differentiate between primary engineering controls and secondary engineering controls. • List the secondary engineering control elements of the cleanroom environment. • Contrast work behaviors in the secondary engineering control contributing to reduction of contamination. • Describe the primary engineering controls used in compounding facilities. • Explain the principles of generating a laminar airflow environment in the primary engineering control.
<p>ACPE #: 0204-0000-19-728-H07-P 0204-0000-19-728-H07-T</p> <p>CE Hours: 1.5 Activity Type: Application</p>	<p>Title: Cleanroom Personnel Basics</p> <p>This activity describes the necessary behaviors and competencies cleanroom staff must master to minimize contamination in cleanroom facilities.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> • Identify sources of contamination within the cleanroom environment. • Distinguish necessary behaviors for cleanroom staff. • Assess personnel for appropriate hand hygiene technique. • Evaluate garbing sequence for non-hazardous and hazardous compounding. • Contrast the components and performance characteristics of garb and gloves for non-hazardous drug and hazardous compounding.

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	<ul style="list-style-type: none"> Describe the testing requirements to assess personnel for appropriate garbing technique. Develop a training and assessment program to ensure mastery of core competencies by aseptic processing personnel.
ACPE #: 0204-0000-19-729-H07-P 0204-0000-19-729-H07-T CE Hours: 3.25 Activity Type: Application	<p>Title: Compounding Materials, Equipment, and Resources for Sterile Product Preparation</p> <p>This activity covers supplies, critical sites, equipment, labels, and references essential for sterile product preparation.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> Recognize basic compounding supplies. Differentiate between the critical sites and non-critical sites on syringes, needles, vials, and bags. Assess a vial’s medication label to determine number of doses and ingredients. Select appropriate compounding materials based on review of medication order. Identify the different types of automated compounder pumps used in sterile product preparation. Choose the appropriate compounding equipment needed for various situations. Interpret several types of patient medication labels. Produce a Master Formulation Record and a Compounding Record. Use tertiary resources to find necessary drug information.
ACPE #: 0204-0000-19-730-H07-P 0204-0000-19-730-H07-T CE Hours: 1.25 Activity Type: Application	<p>Title: Getting Started in Sterile Product Preparation</p> <p>This activity describes dosage forms, small and large volume parenterals, routes of administration, parenteral nutrition, and high alert medications.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> Describe the dosage forms, preparation requirements, and routes of administration. Differentiate between small and large volume parenterals. Recognize appropriate routes of administration for compounded sterile preparations. Explain the components and role of parenteral nutrition. Evaluate parenteral nutrition orders and processes. Propose strategies to mitigate risk of high alert medications.
ACPE #: 0204-0000-19-731-H07-P 0204-0000-19-731-H07-T CE Hours: 2 Activity Type: Application	<p>Title: Stability and Sterility: Assigning Beyond Use Dates</p> <p>This activity discusses the factors that influence the stability and sterility of sterile product preparations and considerations when assigning beyond-use dates.</p>

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	<p>Learning Objectives:</p> <ul style="list-style-type: none"> • List factors that influence beyond-use date assignments for compounded sterile preparations. • Describe physical and chemical compatibility criteria for components. • Apply USP <797> risk categories to assigning a proper beyond-use date for compounded sterile preparations. • Recommend beyond-use date for a final compounded sterile preparation using evidence-based information.
<p>ACPE #: 0204-0000-19-732-H07-P 0204-0000-19-732-H07-T</p> <p>CE Hours: 2.75 Activity Type: Application</p>	<p>Title: Aseptic Techniques for Compounding Non-Hazardous Preparations</p> <p>This activity explains “first air,” aseptic technique in horizontal and vertical airflow, reconstituting powders, and appropriate compounding behaviors to prevent or minimize sharps injuries.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> • Define the concept of “first air.” • Illustrate the location of the direct compounding area in horizontal and vertical airflow. • Describe proper methods for disinfecting critical sites on commonly used sterile components. • Distinguish workflow steps and best practices associated with compounding in a laminar airflow workbench and a compounding aseptic isolator. • Describe techniques for reconstituting sterile powders. • Demonstrate placement of hands to prevent disruption of airflow to critical sites when reconstituting powders and withdrawing diluent or medication from vials. • Restate various strategies used to prevent the potential for coring vial stoppers. • Recommend compounding techniques and behaviors that should be used to prevent and address a sharps injury.
<p>ACPE #: 0204-0000-19-733-H07-P 0204-0000-19-733-H07-T</p> <p>CE Hours: 2.5 Activity Type: Application</p>	<p>Title: Small Volume Parenterals and Other Non-Hazardous Preparations</p> <p>This activity covers techniques and procedures associated with compounding medication cassettes and other “specials” including epidural, intrathecal, and ophthalmic preparations.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> • Describe techniques for compounding medication cassettes. • Practice calculations commonly used with medication cassettes. • Interpret procedures to accurately measure components using principles of volumetric accuracy. • Describe techniques for compounding ‘specials’ including epidural, intrathecal, and ophthalmic preparations.

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<p>ACPE #: 0204-0000-19-734-H07-P 0204-0000-19-734-H07-T</p> <p>CE Hours: 1 Activity Type: Application</p>	<ul style="list-style-type: none"> • Differentiate situations when sterile filtration and/or preservative-free ingredients must be utilized when compounding special administration medications. <p>Title: Nonsterile to Sterile Compounding</p> <p>This activity describes the general requirements and considerations associated with nonsterile to sterile compounding.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> • Explain the requirements under section 503A of the Drug Quality and Safety Act that apply to bulk drug ingredients. • Interpret the requirements under USP <797> for nonsterile to sterile compounding. • Discuss the facility and environmental requirements for nonsterile to sterile compounding. • Evaluate product release requirements for nonsterile to sterile compounding. • Assess the critical quality attributes that must be within limits to guarantee a quality compounded sterile preparation from non-sterile components. • Describe the installation and operational validation process for sterilization equipment. • Distinguish components of quality control.
<p>ACPE #: 0204-0000-19-735-H07-P 0204-0000-19-735-H07-T</p> <p>CE Hours: 2.25 Activity Type: Application</p>	<p>Title: Maintaining the Cleanroom Environment</p> <p>This activity discusses the important elements of cleaning and environmental monitoring required to maintain the cleanroom environment.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> • Identify important considerations for designing a facility cleaning plan. • Describe the process to clean and disinfect classified areas. • Discuss how to build environmental control / monitoring skills into a compounded sterile preparation training program. • Contrast the required supplies and cleaning/disinfecting agents used in laminar airflow systems. • Differentiate cleaning a primary engineering control and a hazardous biological safety cabinet. • Explain how environmental conditions are measured and maintained. • Illustrate how to properly perform volumetric air sampling. • Recommend an appropriate action plan based on personnel and environmental monitoring reports.
<p>ACPE #: 0204-0000-19-736-H07-P</p>	<p>Title: Handling Hazardous Drugs: Part 1</p>

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<p>0204-0000-19-736-H07-T</p> <p>CE Hours: 2 Activity Type: Application</p>	<p>This activity covers the scope and requirements associated with USP chapter <800>, developing Assessments of Risk, and how to evaluate your organization’s current compliance.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> • Describe the key requirements of USP <800> Hazardous Drugs – Handling in Healthcare Settings, including limiting risk to personnel, facility design, and safe work practices. • Explain the scope of USP <800>. • Categorize the handling of hazardous drugs in your organization to determine their eligibility for inclusion in your Assessment of Risk. • Create or review an acknowledgement of risk document. • List questions relevant to your organization after reviewing USP <800>. • Assess your organization's current compliance with USP <800>. • Evaluate the organization's storage and compounding areas. • List the three types of containment primary engineering controls used for compounding hazardous drugs. • List the two types of containment secondary engineering controls used for storage and compounding hazardous drugs. • Analyze your organization's most recent certification report. • Interpret pressure gradients, air flow direction, and air changes per hour.
<p>ACPE #: 0204-0000-19-737-H07-P 0204-0000-19-737-H07-T</p> <p>CE Hours: 2.5 Activity Type: Application</p>	<p>Title: Handling Hazardous Drugs: Part 2</p> <p>This activity describes the key responsibilities of the Designated Person, organizing training materials and checklists, and appropriate hazardous drug work practices.</p> <p>Learning Objectives:</p> <ul style="list-style-type: none"> • Organize your organization's training materials. • List the key responsibilities of the Designated Person. • Differentiate personal protective equipment used in hazardous drug compounding from that used in non-hazardous compounding. • Describe work practice from receiving through compounding. • Design a policy and procedure for handling spills. • Apply appropriate strategies to achieve compliance identified in gap analyses. • Create a checklist that can be used for daily, monthly, and annual monitors for facilities and personnel.
<p>ACPE #: 0204-0000-19-738-H07-P 0204-0000-19-738-H07-T</p> <p>CE Hours: 3.5</p>	<p>Title: Managing the Product until Final Check or Disposal</p> <p>This activity discusses the appropriate movement of drugs and supplies used in sterile product preparation from receipt through final check or disposal.</p>

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<p>Activity Type: Application</p>	<p>Learning Objectives:</p> <ul style="list-style-type: none"> • Illustrate the appropriate movement of drugs and supplies into an ISO classified cleanroom environment. • Analyze storage conditions for drugs and supplies to determine compliance with USP <797> and manufacturer recommendations. • Choose appropriate methods of source ingredient verification prior to compounding sterile products. • Identify appropriate components of physical inspection for source ingredients and final products. • Interpret a Master Formulation Record and Compounding Record as part of the compounded sterile preparation verification process. • Describe when sterility testing and bacterial endotoxin testing is a required quality control in the sterile compounding process. • Apply FDA repackaging guidance to ensure compliance with routine cleanroom practices. • Choose appropriate supplies for packaging and repackaging compounded sterile preparations. • Differentiate between storage, transport, and disposal requirements for hazardous and non-hazardous compounded sterile products. • Analyze storage and transport conditions to ensure compounded sterile product integrity is maintained prior to administration. • Choose appropriate waste disposal stream for pharmaceutical products and supplies used in sterile compounding.

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- Ryan A. Forrey, Pharm.D., M.S., FASHP, Becton, Dickinson and Company (BD): employee
- Kevin N. Hansen, Pharm.D., M.S., BCPS, Baxter International, Inc.: speaker and PharMEDium Services, LLC: speaker
- Patricia C. Kienle, B.S.Pharm., M.P.A., FASHP, Cardinal Health: employee and CriticalPoint, LLC: consultant
- Elaine Strauss, Pharm.D., M.S., WorkingBuildings Group: consultant
- All other planners, presenters, reviewers, and ASHP staff report no financial relationships relevant to this activity.

Methods and CE Requirements

This online activity consists of a combined total of 14 learning modules. Pharmacists and pharmacy technicians are eligible to receive a total of 29.00 hours of continuing education credit by completing all 14 modules within this certificate program.

Participants must participate in the entire activity, complete the evaluation and all required components to claim continuing pharmacy education credit online at ASHP eLearning Portal. Follow the prompts to claim credit and view your statement of credit within 60 days after completing the activity.

Important Note – ACPE 60 Day Deadline:

Per ACPE requirements, CPE credit must be claimed within 60 days of being earned – no exceptions! To verify that you have completed the required steps and to ensure your credits have been reported to CPE Monitor, we encourage you to check your NABP eProfile account to validate your credits were transferred successfully before the ACPE 60-day deadline. After the 60 day deadline, ASHP will no longer be able to award credit for this activity.

System Technical Requirements

System Requirements Courses and learning activities are delivered via your Web browser and Acrobat PDF. Users should have a basic comfort level using a computer and navigating web sites.

View the [minimum technical and system requirements](#) for learning activities.