



KEY CHANGES

The following represents key changes from the currently enforceable version of USP Chapter <797> (last major revision in 2008) to the revised USP Chapter <797> (official as of November 1, 2023). The following are the major changes and are not meant to be an exhaustive list of the entirety of all changes made. Some changes will be reported as direct text excerpts from the respective chapter (notated by quotation marks), while others will be reported as a general comment describing the text or change. *Note: Bolding has been added to the text below for emphasis.*

Category	USP <797>, 2008 ¹	USP <797>, 2023 ²	
01. INTRODUCTION AN	01. INTRODUCTION AND SCOPE		
"The use of technologies, techniques, materials, and procedures other than those described in this chapter "	" not prohibited so long as they have been proven to be equivalent or superior with statistical significance to those described herein."	" not prohibited as long as they are noninferior to those described herein and validated for the intended purpose" (e.g., USP <1223>, <1225>)	
Compounded sterile preparations (CSPs) affected	" irrigations for wounds and body cavities "	" Irrigations for internal body cavities [NOTE—irrigations for the mouth, rectal cavity, and sinus cavity are not required to be sterile .]"	
	" aqueous bronchial and nasal inhalations "	"Nasal dosage forms intended for local application are not required to be sterile."	
Hazardous drugs	Covered within the chapter under section Hazardous Drugs as CSPs Allows preparation of a "low volume of hazardous drugs" outside of a negative pressure space as long as two tiers of containment are used (closed-system transfer device with containment primary engineering control)	Removed from chapter and references to follow USP <800> No longer allows preparation of a low volume of hazardous drugs outside of a negative pressure space.	
Radiopharmaceuticals	Covered within the chapter under section Radiopharmaceuticals as CSPs	Removed from chapter and references to follow USP <825>	
Personnel and settings affected	Largely refers to and addresses only compounding personnel	"Any person entering a sterile compounding area, whether preparing a CSP or not, must meet the requirements in 3. Personal Hygiene and Garbing."	
The designated person(s)	Not addressed	"The compounding facility must designate one or more individuals (i.e., the designated person(s)) to be responsible and accountable for the performance and operation of the facility and personnel in the preparation of CSPs and for performing other functions as described in this chapter." A complete list of the designated person responsibilities has been provided as a separate resource.	

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Administration	Standards do not pertain to the clinical administration of CSPs to patients (e.g., implantation, infusion, inhalation)	"For the purposes of this chapter, 'administration' means the direct application of a sterile product or preparation to a single patient by injecting, infusing, or otherwise providing a sterile product or preparation in its final form."
Immediate-use CSPs	"Administration begins not later than 1 hour following the start of the preparation " Does not involve > 3 commercially manufactured packages of sterile nonhazardous products	"Administration begins within 4 h following the start of preparation." "The preparation involves not more than 3 different sterile products." "Personnel are trained and demonstrate competency in aseptic processes as they related to assigned tasks and the facility's SOPs."
Preparation per approved labeling	Strictly following the manufacturers' approved labeling (product package inserts) is considered a CSP and the requirements of the chapter apply	"Compounding does not include mixing, reconstituting, or other such acts that are performed in accordance with directions contained in approved labeling or supplemental materials provided by the product's manufacturer." "The product is prepared as a single dose for an individual patient" "Approved labeling includes information for the diluent, the resultant strength, the container closure system, and storage time."
Proprietary bag and vial system	Does not mention BUDs other than following the manufacturer's instructions for handling and storing	Docking of the proprietary bag and vial system for future activation This is considered compounding and must be performed in accordance with this chapter (ISO Class 5 environment) BUDs must not be longer than the manufacturer's labeling
CSP microbial categories	 CSP Categories Low Risk Low Risk with 12-h BUD Medium Risk High Risk Factors that determine CSP Category Type of manipulation Complexity and length of preparation If any nonsterile ingredient, component, or equipment is used Number of sterile products and packages Number of transfers into any single container Number of doses being prepared Following proper garbing Exposure to lower than ISO class 5 air and duration 	 Categories Category 1 Category 2 Category 3 Factors that determine CSP Category Primarily based on environment/conditions of where the CSP is compounded Level of garbing Environmental testing and monitoring Frequency of application of a sporicidal Based on BUD assignment "Category 1, Category 2, and Category 3 CSPs can be compounded by using only sterile starting ingredients, or by using some or all nonsterile starting ingredients." One (or more) component is non-sterile: sterility of the compound must be achieved through a sterilization process (e.g., terminal sterilization) and must be maintained if it is subsequently manipulated



Category	USP <797>, 2008 ¹	USP <797>, 2023²		
02. PERSONNEL TRAIN	02. PERSONNEL TRAINING AND EVALUATION			
Who needs to be trained and how often	"Personnel who prepare CSPs shall be trained "	Compounders and those who have direct oversight of compounders		
	How often: • Low- and medium-risk level: at	Initially and at least every 6 or 12 months (depends on the individual)		
	least annually	Personnel who do not compound nor have direct oversight of compounders, but are associated with other tasks		
	High-risk level: semi-annually	(e.g., restock or clean/disinfect the SCA, only compound immediate-use CSPs):		
		Defined by facility SOPs		
Initial garbing competency	Compounders need to pass garbing competency evaluations before	Garbing competency evaluations include:		
evaluations	beginning to prepare CSPs	Visual observation		
		Gloved fingertip and thumb sampling (GFT) of both hands		
		Compounders and those who have direct oversight of compounders		
		" must complete an initial garbing competency evaluation no fewer than 3 separate times. The 3 successful completions must be in succession "		
		Remediation of failed competency		
		" failure of any of the 3 initial garbing competency evaluations requires repeat testing until personnel successfully completes 3 evaluations in a row."		
Ongoing garbing	Visual observation of hand hygiene	Compounders		
competency evaluations	and garbing	Category 1 and 2: at least every 6 months		
	At least annually Claved fing parting and through appending.	Category 3: at least every 3 months		
	Gloved fingertip and thumb sampling	Those who have direct oversight of compounders		
	Low/medium risk – at least annually	At least every 12 months		
	High-risk – at least semiannually			
Initial aseptic	Compounders need to pass media-fill	Aseptic manipulation evaluations include:		
manipulation competency	testing of aseptic manipulation skills before beginning to prepare CSPs	Visual observation		
evaluations		Media-fill testing with post-GFT		
		Surface sampling		
		Compounders and those who have direct oversight of compounders		
		Must complete 1 successful aseptic manipulation competency evaluation		
		Remediation of failed competency		
		"A failure in the media fill, gloved fingertip and thumb sampling, or surface sample constitutes an overall failure of the aseptic manipulation competency."		



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Ongoing aseptic	Each person authorized to compound in a low-risk or medium-risk level	Compounders
manipulation competency	environment:	Category 1 and 2: at least every 6 months
evaluations	At least annually	Category 3: at least every 3 months
	Each person authorized to compound	Those who have direct oversight of compounders:
	in a high-risk level environment:	At least every 12 months
Gloved fingertip and	At least semiannually Incubate sample at 30-35 C for 2-3	"Incubate the media device at 30-35 C for no less than 48 h
thumb sampling incubation standards	days	and then at 20-25 C for no less than 5 additional days."
Media-fill testing incubation standards	Incubate sample at 20-25 C or 30-35 C for 14 days	"Incubate the final containers at 20-25 C and 30-35 C for a minimum of 7 days at each temperature band "
		"The order of the incubation temperatures must be described in the facility's SOPs"
Action levels for gloved fingertip and	0 cfu	After garbing: >0 cfu
thumb sampling		After media-fill testing: >3 cfu
		Action levels based on total cfu count from both hands
03. PERSONAL HYGIEN		
Order of handwashing and garbing	Gave a specific order for garbing and handwashing	Order of handwashing and garbing is determined by the placement of the sink
	Sterile gloves could be donned in the buffer room	Order of garbing must be described by facility's SOPs
	Suiter 100m	"Donning and doffing garb should not occur in the same area at the same time"
		"Sterile gloves must be donned in a classified room or SCA"
Hand hygiene	Allows use of hand dryers	Hand dryers must not be used
	Does not mention soap containers	Disposable soap containers must not be refilled or topped off – need to be replaced
Sanitizing hands	" perform antiseptic hand cleansing with an alcohol-based surgical hand scrub with persistent activity."	Do not need an agent with persistent killing
Reusing garb	Allows gown to be reused if used on the same work day	Category 1 and Category 2
	the sume work day	" gowns may be reused within the same shift by the same person if the gown is maintained in a classified area or adjacent to, or within, the SCA in a manner that prevents contamination."
		Other garb cannot be reused and should be discarded or laundered before reuse
		Category 3
		"Disposable garbing items must not be reused, and laundered garb must not be reused without being laundered and resterilized with a validated cycle."
		"The facility's SOPs must describe disinfection procedures for reusing goggles, respirators, and other reusable equipment."



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Garbing for category 3	Not applicable	"If the facility compounds Category 3 CSPs, additional garbing requirements must be continuously met in the buffer room in which Category 3 CSPs are prepared."
		No exposed skin (I.e., face and neck must be covered)
		All low-lint outer garb must be sterile
		Disposable garb cannot be reused
		Laundered garb cannot be reused until it is laundered and re-sterilized
		Facility's SOPs describe disinfection procedures for reusing goggles, respirators, and other reusable equipment
04. FACILITIES AND EN	GINEERING CONTROLS	
ISO classification of particulate matter	Particle count listed as m³ and ft³	Particle count is only listed as m ³
Use of isolators	PECs shall be located within a restricted access ISO Class 7 buffer area, with exceptions for CAI/CACI which would allow for BUD's	The exception for CAI/CACI's has been removed; to obtain Category 2 CSP BUD's, the CAI/CACI must be placed in an ISO Class 7 buffer room located within a cleanroom suite
	equivalent to a full cleanroom suite in a segregated compounding area when certain conditions are met	Alternatively, a pharmaceutical isolator (different type of engineering control than a CAI/CACI) can be placed in an ISO Class 8 environment without the need for an anteroom
Air exchange requirements	Does not address ISO Class 8 ACPH requirements	ISO Class 8 room: >20 ACPH
Cleanroom	Not addressed	Term to describe ISO-classified anteroom and buffer room
Cleanroom suites: access doors and seals	Not addressed	Seals should not be installed at doors between buffer rooms and anterooms
		Access doors should be hands-free
Precision and accuracy of pressure differentials	Listed as 0.02 (two decimal places), broad	Listed as 0.020 (three decimal places), narrow
Humidity requirements	Does not mention humidity	" should be maintained at a relative humidity of 60% or below "
05. CERTIFICATION AN	D RECERTIFICATION	
Certification of PEC and SEC	"Certification procedures such as those outlined in the CETA Certification Guide for Sterile Compounding Facilities shall be used."	All professional organizations have been removed: " independently certified using the requirements in this chapter and when applicable, manufacturer specifications."
06. MICROBIOLOGICAL	AIR AND SURFACE MONITORING	
Viable air sampling – timing and locations	At least every 6 months for all compounds	 Category 1 and Category 2: At least every 6 months Category 3 Within 30 days before the start of any Category 3
		compounding • At least monthly
		- At least monthly



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Viable air sampling – incubation standards	TSA: • 30-35 C for 48 to 72 h	Incubate at 30-35 C for no less than 48 h then incubate at 20-25 C for no less than 5 additional days
	Fungal media: • 26-30 C for 5 to 7 days	"To shorten overall incubation period, two sampling media devices may be collected for each sample location and incubated concurrently"
		Incubate one at 30-35 C for no less than 48 h and the other at 20-25 C for no less than 5 days
Surface sampling – timing and locations	"Surface sampling shall be performed in all ISO classified areas on a periodic basis"	Locations: • Equipment contained within the PEC • Staging or work area(s) near the PEC • Frequently touched surfaces Category 1 and 2 • At least monthly Category 3 • At least weekly • Prior to assigning a BUD longer than the limits established for Category 2 CSPs
Surface sampling –	Action levels	Action levels
action levels	• ISO Class 5: >3	• ISO Class 5: >3
	• ISO Class 7: >5	• ISO Class 7: >5
	• ISO Class 8 or worse: >100	• ISO Class 8: >50
Identifying microorganisms and Corrective Actions	Identification of microorganisms (at least the genus level) is required regardless of cfu count	If action levels specified for air and surface sampling are exceeded, " an attempt must be made to identify any microorganism recovered to the genus level"
	Mention of highly pathogenic	Does not mention highly pathogenic microorganisms
	microorganisms (e.g., gram-negative rods, coagulase <i>Staphylococcus</i> , molds and yeasts) must be	"The extent of the investigation should be consistent with the deviation and should include an evaluation of trends"
	immediately remedied regardless of cfu count	"Data collected in response to corrective actions must be reviewed to confirm that the actions taken have been effective."
		"The corrective action plan must be dependent on the cfu count and the microorganism recovered."
07. CLEANING, DISINFECTING, AND APPLYING SPORICIDAL DISINFECTANTS AND STERILE 70% IPA		
Minimum frequency for cleaning and disinfecting surfaces	Does not split up minimum frequency based on method (e.g., cleaning, disinfecting)	Minimum frequency for cleaning is broken down by cleaning, disinfecting, and applying sporicidal disinfectant



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Cleaning/disinfecting supplies	Does not specify the type of material	Cleaning and disinfecting supplies (e.g., wipers, sponges, pads, mop heads)
		Must be low-lint
		Should be disposable
		Reusable cleaning tools must be dedicated for use and not be removed from classified areas or SCA and be made of cleanable materials (e.g., not wood or any other porous material)
		"Cleaning, disinfecting and sporicidal agents used within the PEC must be sterile ." Sterile water must be used when diluting concentrated agents for use in the PEC.
08. INTRODUCING ITEM	IS INTO THE SEC AND PEC	
No major changes		
09. EQUIPMENT, SUPPL	IES, AND COMPONENTS	
No major changes		
10. STERILIZATION AND	DEPYROGENATION	
Biological indicators	Steam Heat - Bacillus stearothermophilus	Steam Heat - Geobacillus stearothermophilus
	, i	Dry Heat – Bacillus atrophaeus
11 MASTED FORMULAT	Dry Heat – Bacillus subtilis ON AND COMPOUNDING RECORDS	
		Must be created for all CSPs prepared for more than
Master formulation records (MFR)	Specific requirements not listed	Must be created for all CSPs prepared for more than one patient or when using non-sterile components
		Any changes or alterations must be approved and documented based on facility's SOPs
		Requirements for MFR are listed out in section
Compounding records (CR)	Specific requirements not listed	Must be created for all Category 1, Category 2, and Category 3 CSPs and for immediate-use CSPs when prepared for more than one patient
10 DELEACE MADE CELO		Requirements for CR are listed out in section
12. RELEASE INSPECTION		
Maximum batch size	Not addressed	"The maximum batch size for all CSPs requiring sterility testing must be limited to 250 final yield units."
Sterility testing	"A method not described in the <i>USP</i> may be used if verification results	Specifies a <i>USP</i> chapter " or a validated alternative method (see <1223>) that is
	demonstrate that the alternative is at least as effective and reliable "	noninferior to <71> testing."
Number of CSPs needed to send for sterility testing	Does not specify number of CSPs needed to be sent for sterility testing	Number of CSPs sent for sterility testing depends on number of CSPs to be compounded in a single batch 1-39 CSPs – must send 10% of the number of CSPs
		prepared, rounded up to the next whole number
		 >40 CSPs – must use sample sizes specified in <71>, Table 3



Category	USP <797>, 2008 ¹	USP <797>, 2023 ²
Sterility testing requirements 13. LABELING	Required for high-risk level CSPs under certain circumstances: • >25 identical individual singledose packages • Multiple-dose vials for administration to multiple patients • Exposed longer than 12 h at 2-8 C and longer than 6 h at warmer than 8 C before they are sterilized	Category 1 – not required Category 2 – based on BUD Category 3 – required
Compounding notification on label	Not addressed	"The labeling on the CSP should indicate that the preparation is compounded."
14. ESTABLISHING BEYO	OND-USE DATES	
Establishing a BUD for a CSP	Factors that determine a BUD for risk categories • Storage conditions • Information gathered from professional sources (e.g., sterility studies)	 Factors that determine Category 1 BUDs Storage conditions (e.g., controlled room temperature, refrigerator) Factors that determine Category 2 BUDs Compounding method (e.g., aseptic process, terminally sterilized) If sterility testing is performed Starting component of compound (e.g., sterile, nonsterile) Storage conditions Additional requirements needed for longer BUDs in Category 3 CSPs for: Increase use of sporicidal disinfectants Increase of environmental monitoring Use of sterile garb Stability determination Personnel qualification
Non-preserved topical ophthalmic CSPs	Not addressed	"The beyond-use-date of a multiple-dose, aqueous, non-preserved CSP intended for topical, including topical ophthalmic, administration may be assigned in accordance with 14.5 Multiple-Dose CSPs." Requirement for passing antimicrobial effectiveness testing in accordance with <51> is not required only if the preparation is: • Prepared as a Category 2 or Category 3 CSP • For use by a single patient • Labeled to indicate that once opened, it must be discarded after 24 h stored at controlled room temp or 72 h stored under refrigeration



Category	USP <797>, 2008 ¹	USP <797>, 2023 ²		
15. USE OF CONVENTIO	15. USE OF CONVENTIONALLY MANUFACTURED PRODUCTS AS COMPONENTS			
Use of conventionally manufactured single-dose containers	"Single-dose vials exposed to ISO Class 5 or cleaner may be used up to 6 h after initial needle puncture."	"If a single-dose vial is entered or punctured only in an ISO Class 5 or cleaner air, it may be used up to 12 h after initial entry or puncture as long as the labeled storage requirements during that 12-h period are maintained."		
Use of conventionally manufactured pharmacy bulk package	Not addressed	"The pharmacy bulk package must be used according to the manufacturer's labeling (see <659>, General Definitions, Injection Packaging Systems). The pharmacy bulk package must be entered or punctured only in an ISO Class 5 PEC."		
16. USE OF CSPS AS CO	MPONENTS			
Use of compounded multiple-dose CSPs	Not addressed	 When used as a component to compound additional CSPs Required to meet criteria for antimicrobial effectiveness testing and requirements in 14.5 Must be stored in conditions the BUD is based (e.g., refrigerator) After punctured, must not be used longer than assigned BUD or 28 days, whichever is shorter. Remainder must be discarded 		
Use of compounded single-dose CSPs and CSP stock solutions	Not addressed	 When used as a component to compound additional CSPs Must be entered or punctured in ISO Class 5 or cleaner air Must be stored in conditions the BUD is based (e.g., refrigerator) May be used for sterile compounding up to 12 h or its assigned BUD, whichever is shorter. Remainder must be discarded 		
17. SOPS				
Who needs training based on facilities SOPs	Not addressed	"All personnel who perform or oversee compounding or support activities must be trained in the SOPs"		
18. QUALITY ASSURANCE	CE AND QUALITY CONTROL			
Notification and recall of CSPs with out-of- specification limits	Not addressed except for notifying the patient and physician of potential risk	 SOP for recall of out-of-specification limits must contain procedures To determine severity of problem and urgency for implementation and completion of the recall To determine distribution of any affected CSP To identify patients who received the CSP For disposal and documentation of recalled CSP To investigate and document reason for failure 		
Redispensed CSPs	Unopened, unused, returned CSPs may be redispensed when certain conditions are met to ensure the CSP is sterile, pure, and stable	Not specifically addressed; however does not prohibit this practice. Would need to refer to state board of pharmacy for guidance.		
19. CSP HANDLING, STO	DRAGE, PACKAGING, SHIPPING, AND TR	RANSPORT		
No major changes				



Category	USP <797>, 2008 ¹	USP <797>, 2023 ²
20. DOCUMENTATION		
No major changes		
21. COMPOUNDING ALL	ERGENIC EXTRACTS	
Compounding allergenic extract prescription sets	No mention of training or competency evaluation needed for compounders making allergenic extracts	 Requirements for personnel who prepare allergenic extracts Training must be done initially prior to compounding independently and annually Gloved fingertip and thumb sampling on both hands no fewer than 3 separate times needs to be done prior to compounding independently and at least every 12 months Sterile technique of compounders needs to be evaluated at least every 12 months Personnel that have not compounded in 6 months need to be evaluated in all core competencies before resuming their duties

References

- 1. United States Pharmacopeial Convention. General chapter <797> pharmaceutical compounding—nonsterile preparations. USP43-NF38. Rockville, MD: U.S. Pharmacopeial Convention; 2019.
- 2. United States Pharmacopeial Convention. General chapter <797> pharmaceutical compounding—sterile preparations. USP-NF 2023, Issue 1, November 1, 2022, official as of November 1, 2023.

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