

California State Board of Pharmacy 2720 Gateway Oaks Drive, Ste. 100 Sacramento, CA 95833 Phone: (916) 518-3100 Fax: (916) 574-8618

Business, Consumer Services and Housing Agency
Department of Consumer Affairs
Gavin Newsom, Governor



Yellow Highlights reflect the changes for Legislation for 2025.

www.pharmacy.ca.gov

2024 Legislation shown by strikethrough for deleted language and dashed underline for added language.

HOSPITAL PHARMACY SELF-ASSESSMENT

Title 16 of the California Code of Regulations section 1715 requires the pharmacist-in-charge of each pharmacy licensed under section 4037 or 4029 of the Business and Professions Code (BPC) to complete a self-assessment of the pharmacy's compliance with federal and state pharmacy law. The assessment shall be performed before July 1 of every odd-numbered year. The pharmacist-in-charge must also complete a self-assessment within 30 days whenever (1) a new pharmacy permit has been issued, or (2) there is a change in the pharmacist-in-charge. The primary purpose of the self-assessment is to promote compliance through self-examination and education.

The self-assessment must be completed in its entirety. It may be completed online, printed, initialed, signed, and readily available in the pharmacy. Signatures and initials shall be an original handwritten signature or initial on the self-assessment form. Do not copy a previous assessment.

Notes: If dispensing prescriptions for outpatient use, a Community Pharmacy Self-Assessment/Hospital Outpatient Pharmacy Self-Assessment (17M-13, pursuant to 16 CCR 1715) must be completed also. A hospital that compounds drug products must also complete the Compounding Self-Assessment (17M-39 pursuant to 16 CCR 1735.2(k).)

Each self-assessment must be kept on file in the pharmacy for three years after it is performed.

Pharmacy Name:		
Address:	Pho	one:
Ownership: Sole Owner For Sole Owner Non-Licensed Owner	Partnership □ Corporat ner □ Other (please s	ion □ LLC □ Trust pecify)
License #: Exp. Date:	Other Licens	pecify) se #: Exp. Date:
Licensed Sterile Compounding Lice	ense # E	xpiration:
Accredited by (optional):	From:	To:
Centralized Hospital Packaging #: Exp		Exp. Date:
DEA Registration #:	Exp. Date:	Date of DEA Inventory:
Hours: Weekdays Sat	Sun	24 Hours

PIC:	RPH #	Exp. Date:
	icists, interns, technicians): Pharmacist, DEA = Drug Enforceme	nt Administration.
	RPH#	Exp. Date:
	APH #	Exp. Date:
	DEA #	Exp. Date:
	RPH#	Exp. Date:
	APH #	Exp. Date:
	DEA #	Exp. Date:
	RPH#	Exp. Date:
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3	TCH#	Exp. Date:

HOSPITAL PHARMACY SELF-ASSESSMENT

All references to the California Code of Regulations (CCR) are Title 16 unless otherwise noted.

Please mark the appropriate box for each question. If "NO," enter an explanation on "CORRECTIVE ACTION or ACTION PLAN" lines below. If more space is needed, you may add additional sheets.

1. Ph Yes No N/A	narmacy
	1.1. The pharmacy is secure and has provisions for effective control against the theft of dangerous drugs and devices. (BPC 4116, 4117, CCR 1714)
	1.2. The pharmacy has procedures in place to take action to protect the public when a licensed individual employed by or with the pharmacy is discovered or known to be chemically, mentally, or physically impaired to the extent it affects their ability to practice the profession or occupation authorized by their license, or is discovered or known to have engaged in the theft, diversion, or self-use of dangerous drugs(BPC 4104[a])
	1.3. The pharmacy has written policies and procedures for addressing chemical, mental, or physical impairment, as well as theft, diversion, or self-use of dangerous drugs, among licensed individual employed by or with the pharmacy. (BPC 4104[b])
	1.4. The pharmacy reports to the board within 14 days of the receipt or development of the following information with regard to any licensed individual employed by or with the pharmacy: (1) any admission by a licensed individual of chemical, mental, or physical impairment affecting their ability to practice; (2) Any admission by a licensed individual of theft, diversion, or self-use of dangerous drugs; (3) Any video or documentary evidence demonstrating chemical, mental, or physical impairment of a licensed individual to the extent it affects their ability to practice; (4) Any video or documentary evidence demonstrating theft, diversion, or self-use of dangerous drugs by a licensed individual; (5) Any termination based on chemical, mental, or physical impairment of a licensed individual to the extent it affects their ability to practice; (6) Any termination of a licensed individual based on theft, diversion, or self-use of dangerous drugs. (BPC 4104[c])
	1.5. The pharmacy maintains a supply of medications which are accessible without entering the pharmacy during hours when the pharmacy is closed, and the pharmacist is not available. Access is limited to designated registered nurses. (22 CCR 70263[n])
	1.6. The pharmacy is of sufficient size and has an unobstructed area to accommodate the safe practice of pharmacy. (CCR 1714[b])
	1.7. The pharmacy premises, fixtures, and equipment are maintained in a clean and orderly condition. (CCR 1714[c])
	1.8. The pharmacy sink has hot and cold running water. (CCR 1714[c])
	1.9. The pharmacy has a readily accessible restroom. (CCR 1714[g])
	1.10. The original board-issued pharmacy license and the current renewal are posted where they may be clearly read by the purchasing public. (BPC 4032, 4058)

Yes No N/A	1.11. Pharmacists, interns, pharmacy technicians, and pharmacy technician trainees wear nametags, in 18-point type, that contain their name and license status. (BPC 680, 4115.5[e], CCR 1793.7[c])
	1.12. Does the pharmacy compound sterile drugs?
	(If yes, complete the Compounding Self-Assessment required by CCR 1735.2[k])
	1.13. The pharmacy is subscribed to the board's email notifications. (BPC 4013)
	Date Last Notification Received:
	Email address registered with the board:
	1.14. For a pharmacy whose owner owns two or more pharmacies, the pharmacy receives the board's email notifications through the owner's electronic notice system. (BPC 4013[c])
	Date Last Notification Received:
	Email address registered with the board:
	1.15. All medicinal cannabis is stored in a locked container in the patient's room, other designated areas, or with the patient's primary caregiver and is retrieved, administered, handled, removed and disposed in accordance with HSC 1649.1, 1649.2, 1649.3, 1649.4.
CORREC	TIVE ACTION OR ACTION PLAN:
	sing Stations
Yes No N/A	2.1. Adequate space is available at ward or nursing station for the storage of drugs and preparation of medication dosesAll such spaces and areas can be locked and are accessible to authorized personnel only. (22 CCR 70269)
	2.2. The pharmacist, intern pharmacist, or pharmacy technician completes the monthly inspections of all floor stock and drugs maintained in nursing stations. Any irregularities are reported to the director of nursing services, and as required by hospital policy. (BPC 4119.7[c], 4115[j], 22 CCR 70263[q][10])
	 2.2.1. An intern pharmacist shall report any irregularities to the pharmacist. (BPC 4119.7[c])
	 2.2.2. A pharmacy technician shall report any irregularities to the pharmacist-in- charge and to the director or <u>chief executive officer</u> of the health care facility within 24 hours. (BPC 4115[ji][3])
CORREC	TIVE ACTION OR ACTION PLAN:

3. Delivery of Drugs

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Yes No N/A	.1. Delivery to the pharmacy of dangerous drugs and dangerous devices are only delivered to the licensed premise and signed for and received by a pharmacist. (BPC 4059.5[a])		
	3.2. Deliveries to a hospital pharmacy may be made to a central receiving location within the hospital. However, the dangerous drugs or dangerous devices shall be delivered to the licensed pharmacy premise within one working day following receipt by the hospital, and the pharmacist on duty at that time shall immediately inventory the drugs or devices. (BPC 4059.5[c])		
	.3. A pharmacy may take delivery of dangerous drugs and dangerous devices when the pharmacy is closed and no pharmacist is on duty if all of the following requirements are met: (BPC 4059.5[f])		
	□ 3.3.1. The drugs are placed in a secure storage facility in the same building as the pharmacy; (BPC 4059.5[f][1])		
	 3.3.2. Only the pharmacist-in-charge or a pharmacist designated by the pharmacist in-charge has access to the secure storage facility after dangerous drugs or dangerous devices have been delivered; (BPC 4059.5[f][2]) 		
	 3.3.3. The secure storage facility has a means of indicating whether it has been entered after dangerous drugs or dangerous devices have been delivered; (BPC 4059.5[f][3]) 		
	 3.3.4. The pharmacy maintains written policies and procedures for the delivery of dangerous drugs and dangerous devices to a secure storage facility; and (BPC 4059.5[f][4]) 		
	□ 3.3.5. The agent delivering dangerous drugs and dangerous devices pursuant to the subdivision leaves documents indicating the name and amount of each dangerous drug or dangerous device delivered in the secure storage facility. The pharmacy shall be responsible for the dangerous drugs and dangerous devices delivered to the secure storage facility. The pharmacy shall also be responsible for obtaining and maintaining records relating to the delivery of dangerous drugs and dangerous devices to a secure storage facility. (BPC 4059.5[f][5])		
□□□ 3.4	Prior to, or at the time of, accepting ownership of a product included in the Drug Supply Chain Security Act from an authorized trading partner, the pharmacy is provided transaction history, transaction information, and a transaction statement. (21 USC 360eee-1[d][1][A][i])		
□□□ 3.9	Prior to, or at the time of, each transaction in which the pharmacy transfers ownership a product included in the Drug Supply Chain Security Act to an authorized trading partner, the subsequent owner is provided transaction history, transaction information, and a transaction statement for the product. Note: This requirement does not apply to sales by a pharmacy to another pharmacy to fulfill a specific patient need. (21 USC 360eee-1[d][1][A][ii])		

Yes No N/A	5. The pharmacy captures transaction information (including lot level information, if provided), transaction history, and transaction statements, as necessary to investigate a suspect product, and maintains such information, history, and statements for not less than 6 years after the transaction. (21 USC 360eee-1[d][1][A][iii])
□□□ 3.7	. The pharmacy is aware, effective November 27, 2020, pharmacies are required by the Drug Quality and Security Act (DQSA), to have pharmacy lot-level traceability and by November 27, 2023 unit-level traceability. The pharmacy has lot-level and unit-level traceability in accordance with the Drug Quality and Security Act (DQSA). (21 USC 360eee-1[d][2] and 582[g][1])
CORREC	TIVE ACTION OR ACTION PLAN:
4. Dru	g Stock
Yes No N/A	4.1. The drug stock is clean, orderly, properly stored, properly labeled and in-date. (21 USC sections 331, 351, 352, BPC 4169[a][2]-[4], 4342, HSC 111255, 111335, CCR 1714 ([b]), 22 CCR 70263[q])
	4.2. All ward/floor drug stock and drug supplies that are maintained for access when the pharmacist is not available are properly labeled and stored. Records of drugs taken from the drug stock or drug supplies must be maintained and the pharmacist must be notified. (22 CCR 70263[n])
	4.3. Preferentially priced drugs are furnished solely or predominately to inpatients in accordance with provisions of the Nonprofit Institutions Act (15 USC 13c). Such drugs also may be dispensed pursuant to prescriptions for inpatients at the time of discharge, for employees of the hospital, or on an emergency basis for a walk-in customer (provided that sales to walk-ins do not exceed one percent of the pharmacy's total prescription sales) or to any person in the occasional emergency situation where no other sources are readily available in the community to meet the emergency need. (BPC 4380, CCR 1710[a])
	4.4. All unit-dose drugs received from a centralized hospital packaging pharmacy are correctly labeled, are barcoded, and the barcode is readable at the patient's bedside. (BPC 4128.4, 4128.5)
	4.5. All drugs are maintained in accordance with national standards regarding the storage area and refrigerator or freezer temperature and manufacturer's guidelines. (BPC 4119.7[b]
	4.6. Dangerous drugs or dangerous devices are purchased, traded, sold or transferred with an entity licensed with the board as a wholesaler, third-party logistics provider, pharmacy or a manufacturer, and provided the dangerous drugs and devices: (BPC 4059.5, 4169, CCR 1718.1)
	4.6.1. Are not known or reasonably should not be known to the pharmacy as being adulterated.

	4.6.2. Are not known or reasonably should not be known to the pharmacy as being misbranded.		
	□ 4.6.3. Are not expired.		
Yes No N/A	4.7. If the pharmacy has reasonable cause to believe a dangerous drug or dangerous device in, or having been in its possession is counterfeit or the subject of a fraudulent transaction, the pharmacy will notify the board within 72 hours of obtaining that knowledge. (BPC 4107.5)		
	4.8. The pharmacy does not furnish dangerous drugs or dangerous devices to an unauthorized person. (BPC 4163)		
	4.9. An automated unit dose system (AUDS) operated by a licensed hospital pharmacy as defined by BPC 4029, and used solely to provide doses administered to patients while in a licensed general acute care hospital facility shall be exempted from the requirement of obtaining an ADDS license if the hospital pharmacy owns or leases the AUDS and owns the dangerous drugs or devices in the AUDS. The AUDS shall comply with all other requirements for an ADDS (i.e. Security, Record keeping, Self-Assessment, Quality Assurance, etc.) and maintain a list of the location of each AUDS it operates. (BPC 4119.11[b][3], 4427.2, 4427.65)		
CORRE	CTIVE ACTION OR ACTION PLAN:		
	armacies That Donate Drugs to a Voluntary County-Approved Drug Repository and tribution Program 5.1. Does the pharmacy donate to or operate a county-approved Voluntary Drug Penesitory and Distribution Program?		
(If yes c	Repository and Distribution Program? omplete Section 30 [donate drugs] or Section 31 [operate program] of this Self-Assessment.)		
(11 yes, e	□ 5.1.1. The pharmacy that donates medications to or operates a voluntary county approved drug repository and distribution program meets all the requirements as specified in law. (HSC 150200, 150201, 150202, 150202.5, 150203, 150204, 150204.5, 150204.6, 150205, BPC 4169.5)		
	5.1. The hospital pharmacy donates medications to a county-approved drug repository and distribution program, and meets the following requirements: (HSC 150202, 150202.5, 150204)		
	∃ 5.1.1. The hospital pharmacy is licensed by and is not on probation with the		
	California State Board of Pharmacy, and (HSC 150202.5)		
	California State Board of Pharmacy, and (HSC 150202.5) 5.1.2. The hospital pharmacy's primary or sole type of pharmacy practice is limited to skilled nursing facility, home health care, board and care, or mail order.		

	5.3. Drugs that are donated are unused, unexpired and meet the following requirements: (HSC 150202.5, 150204[c])
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	5.4. The hospital pharmacy follows the same procedural drug pedigree requirements for donated drugs as it does for drugs purchased from a wholesaler or directly from a drug manufacturer. (HSC 150204[n])
CORREC	TIVE ACTION OR ACTION PLAN:
CONNEC	
	nacist-in-Charge (PIC)
	macist-in-Charge (PIC) 6.1. The pharmacy has a PIC who is responsible for the daily operation of the pharmacy. (BPC 4101, 4113, 4305, 4330, CCR 1709, 1709.1)
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6. Pharm Yes No N/A □□□□	6.1. The pharmacy has a PIC who is responsible for the daily operation of the pharmacy. (BPC 4101, 4113, 4305, 4330, CCR 1709, 1709.1)6.2. The PIC has adequate authority to assure the pharmacy's compliance with laws
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<u> </u>	6.7. The PIC or pharmacist on duty shall immediately notify the store management of any conditions that present an immediate risk of death, illness, or irreparable harm to patients, personnel, or pharmacy staff. If the conditions are not resolved within 24 hours, the PIC or pharmacist on duty shall ensure the board is timely notified. (BPC 4113[d][1]
CORRE	CTIVE ACTION OR ACTION PLAN:
7. Dutie	es of a Pharmacist
Yes No N/A	7.1. A pharmacist: (BPC 4019, 4051, 4052, 4052.2, CCR 1717[c], CCR 1793.1, CCR 1793.7)
	 7.1.1. Receives a chart order for an inpatient; (BPC 4019, 4051-[b], 4052, 4052.2, CCR 1717, CCR 1793.1[a])
	 7.1.2. Identifies, evaluates and interprets the chart order; (CCR 1717[c], CCR 1793.1[c])
	 7.1.3. Reviews patient's drug regimen and interprets the clinical data in the patient's medication record; (BPC 4052.1[a][4], 4052.2[a][4], CCR 1793.1[d])
	 7.1.4. Consults with any prescriber, nurse or health care professional; (CCR 1793.1[e])
	□ 7.1.5. Calculates drug doses; (BPC 4052-[a][3], 4052.2-[a][3], 4052.2-[a][4])
	 7.1.6. Supervises the packaging of drugs and checks the packaging procedures and products upon completion; (CCR 1793.1[f])
	7.1.7. Is responsible for all activities of pharmacy technicians, interns and clerks related to the furnishing of drugs to ensure that all such activities are performed completely, safely and without risk of harm to patients; (CCR 1793.7[e])
	7.1.8. Performs any other duty which federal or state law or regulation authorizes only a registered pharmacist to perform; and performs all functions which require professional judgment. (BPC 4052, 4052.2, CCR 1793.1[g])
	7.2. Pharmacists in a licensed health care facility who are performing the following functions are doing so in accordance with the hospital's policies, procedures and protocols which have been developed by health professionals including physicians, pharmacists, and registered nurses, with the concurrence of the facility administrator: (BPC 4027, 4051, 4052, 4052.2)
	 7.2.1. Ordering or performing routine drug therapy-related patient assessment procedures; (BPC 4052.1[a][1]; 4052.2[a][1])
	 7.2.2. Ordering drug therapy-related laboratory tests; administering drugs or biologicals by injection; (BPC 4052.1[a][2],-[3]; 4052.2[a][2],-[3])
	 7.2.3. Initiating or adjusting the drug regimen of a patient; (BPC 4052.1[a][4], BPC 4052.2[a][4])

	these functions, the pharmacist must have either (1) successfully completed clinical residency training, or (2) demonstrated clinical experience in direct patient care delivery as specified in BPC section 4052.2[d]. (BPC 4052.4)		
	7.2.5. A pharmacist may perform any aspect of any FDA-approved or authorized test that is classified as waived pursuant to the federal Clinical Laboratory Improvement Amendments of 1988 (42 USC Sec 263a) and the pharmacist completes the testing in a pharmacy laboratory that is licensed in California as a laboratory pursuant to BPC section 1265 unless otherwise authorized in law. The pharmacist has completed necessary training as specified in the pharmacy's policies and procedure maintained in subsection be of BPC section 4119.10 and that allows the pharmacist to demonstrate sufficient knowledge of the illness, condition or disease being tested as applicable. (BPC 4052.4)		
Yes No N/A □□□□	7.3. The pharmacist who is authorized to issue an order to initiate or adjust a controlled substance therapy is personally registered with the federal Drug Enforcement Administration. (BPC 4052[b])		
	7.4. All pharmacists have submitted an application to the Department of Justice to obtain approval to access information online regarding the controlled substance history of a patient. Upon approval, the DOJ shall release to the pharmacist or their delegate the CURES information for an individual under the pharmacist's care. (HSC 11165.1)		
	7.5. All pharmacists have joined the board's email notification list. (BPC 4013)		
	7.6 The hospital pharmacist (or pharmacy technician or an intern pharmacist if both requirements of BPC 4118.5(b) are met) shall obtain an accurate medication profile or list for each high-risk patient upon admission of the high-risk patients if the hospital has more than 100 beds, the accurate medication profile is acquired during hospital pharmacy's hours of operation. (BPC 4118.5)		
	7.7. The pharmacist may initiate, adjust or discontinue drug therapy for a patient under a collaborative practice agreement with any health care provider with prescriptive authority. The collaborative practice agreement may be between a single or multiple pharmacists and a single or multiple health care providers with prescriptive authority. (BPC 4052[a][13], [14])		
	7.8. Only a prescriber, a prescriber's authorized agent, or a pharmacist may electronically enter a prescription or an order, as defined in BPC 4019, into a pharmacy's or hospital's computer from any location outside of the pharmacy or hospital with the permission of the pharmacy or hospital. (BPC 4071.1)		
	7.9. A pharmacist located and licensed in the state may, on behalf of a health care facility licensed pursuant to Chapter 2 (commencing with Section 1250) of Division 2 of the Health and Safety Code, from a location outside the facility, verify medication chart orders for appropriateness before administration consistent with federal requirements, as established in the health care facility's policies and procedures. The health care facility shall maintain a record of the pharmacist's verification of the medication chart order that meets the same requirements as those described in Sections 4081 and 4105. (BPC 4071.1[d][1], 4071.1[d][2])		

CORRECT	IVE A	CTION OR ACTION PLAN:
8. Duties	s of a	n Advanced Practice Pharmacist
Yes No N/A		The advanced practice pharmacist has received an advanced practice pharmacist nse from the board and may do the following: (BPC 4016.5, 4210)
		8.1.1 Perform patient assessments, order and interpret drug therapy-related tests, and refer patients to other health care providers; (BPC 4052.6[a])
		8.1.2 Participate in the evaluation and management of diseases and health conditions in collaboration with other health care providers; (BPC 4052.6[a])
		8.1.3 Initiate, adjust or discontinue drug therapy and shall promptly transmit writter notification to, or enter the appropriate information into a patient record system shared with the patient's primary care provider or diagnosing provider; (BPC 4052.6[a][5], [b])
		8.1.4 Adjust or discontinue drug therapy and promptly transmit written notification to the patient's diagnosing prescriber or enters the appropriate information in a patient's record system shared with the prescriber; (BPC 4052.6[b])
		8.1.5 Prior to initiating or adjusting a controlled substance therapy, the advance practice pharmacist is personally registered with the federal Drug Enforcement Administration; (BPC 4052.6[d])
		8.1.6 Ordering of tests is done in coordination with the patient's primary care provider or diagnosing prescriber, including promptly transmitting written notification to the prescriber or entering information in a patient record system shared with the prescriber. (BPC 4052.6[e])
CORRECT	IVE A	CTION OR ACTION PLAN:
Yes No N/A	.1. Int	Intern Pharmacist ern pharmacists are performing all the functions of a pharmacist only under the ect supervision of a pharmacist, and the pharmacist is supervising no more than two erns at any one time. (BPC 4023.5, 4030, 4114, 4119.6, 4119.7, CCR 1726) 9.1.1 Stock, replenish and inspect the emergency pharmaceutical supply container and the emergency medical system supplies. (BPC 4119.6) 9.1.2. Inspect the drugs maintained in the health care facility at least once per month. (BPC 4119.7[c])

PIC

	9.2. All prescriptions filled or refilled by an intern are initialed or documented by secure computer entry by a pharmacist prior to dispensing. (CCR 1712[a], 1717[b][1])
	9.3. During a temporary absence of a pharmacist for a meal period or duty-free break, an intern pharmacist does not perform any discretionary duties or act as a pharmacist. (CCR 1714.1[d])
	9.4. The intern hours affidavits are signed by the pharmacist under whom the experience was earned or by the pharmacist-in-charge at the pharmacy while the intern pharmacist obtained the experience, when applicable. (BPC 4209[b], [c], [d]; CCR 1726)
	9.5. All intern pharmacists have joined the board's email notification list. (BPC 4013)
CORREC	TIVE ACTION OR ACTION PLAN:
10. Dutie	es of a Pharmacy Technician
Yes No N/A	of a Final made Footimeran
	10.1. Registered pharmacy technicians are performing packaging, manipulative, repetitive, or other nondiscretionary tasks related to the furnishing of drugs, while assisting and under the direct supervision and control of a pharmacist. The pharmacist is responsible for the duties performed by the pharmacy technician under the pharmacist's supervision. (BPC 4023.5, 4038, 4115, CCR 1793.2, CCR 1793.7)
	10.2. The ratio is not less than one pharmacist on duty for two technicians when filling prescriptions for an inpatient of a licensed health facility. (BPC 4115[gf], CCR 1793.7[f])
	10.3. When prescriptions are dispensed to discharge patients with only one pharmacist, there is no more than one technician <u>performing packaging</u> , <u>manipulative</u> , <u>repetitive</u> , <u>or other nondiscretionary tasks</u> . If a pharmacy technician, under the direct supervision and control of the pharmacist, prepares and administers influenza and COVID-19 vaccines via injection or intranasally, prepares and administers epinephrine, performs specimen collection for tests that are classified as waived under CLIA, receives prescription transfers, and accepts clarification on prescriptions, a second pharmacy technician shall be assisting a pharmacist with performing the tasks as defined in BPC 4115(a). The ratio of pharmacy technicians performing those tasks for additional pharmacists does not exceed 2:1. (BPC 4038, 4115[gf][1], CCR 1793.7[f])
Yes No N/A	10.4. Any function performed by a technician in connection with the dispensing of a prescription or chart order, including repackaging from bulk and storage of pharmaceuticals is verified and documented in writing by a pharmacist or documented by a pharmacist using secure computer entry. (CCR 1712, 1793.7)
	10.5. A pharmacy technician or pharmacy technician trainee wears identification, in 18-point type that identifies them as a pharmacy technician or pharmacy technician trainee (BPC 680, BPC 4115.5[e], CCR 1793.7[d])
	10.6. The pharmacy has a job description for the pharmacy technician and written policies and procedures to ensure compliance with the technician requirements. (CCR 1793.7)

	pharn but m techn	ring a temporary absence of a pharmacist for a meal period or duty-free break, a macy technician may, at the discretion of the pharmacist, remain in the pharmacy may only perform nondiscretionary tasks. Any task performed by the pharmacy nician during the pharmacist's temporary absence is reviewed by the pharmacist. 4115[hg], CCR 1714.1[c])
	allows	e general acute-care hospital has an ongoing clinical pharmacy program and s specially trained pharmacy technicians to check the work of other pharmacy licians when the following conditions are met: (CCR 1793.8)
		10.8.1. Pharmacists are deployed to the inpatient care setting to provide clinical services. 10.8.2. Compounded or repackaged products are previously checked by a pharmacist, then used by the technician to fill unit dose distribution and floor and
		ward stock. 10.8.3. The overall operations are the responsibility of the pharmacist-in-charge. 10.8.4. The pharmacy technician checking technician program is under the direct supervision of the Pharmacist as specified in the policies and procedures.
	a	10.8.5. There is an ongoing evaluation of the program that uses specialized and advanced trained pharmacy technicians to check the work of other pharmacy technicians.
Yes No N/A	10.9. Pha	armacy technician duties include the following:
		10.9.1. Package emergency supplies for use in the health care facility and the hospital's emergency medical system. (BPC 4119, 4115[jɨ])
		10.9.2. Seal emergency containers for use in the health care facility. (BPC 4115[jɨ])
	(10.9.3. Perform monthly checks of the drug supplies stored throughout the health care facility and report any irregularities within 24 hours to the pharmacist-incharge and to the director or chief executive officer. (BPC 4115[jɨ])
Yes No N/A	10.10. Al	Il pharmacy technicians have joined the board's email notification list. (BPC 4013)
Yes No N/A	<u>board</u>	person shall not act as a pharmacy technician without first being licensed by the day a pharmacy technician. A pharmacy technician certification only is not ralent to being licensed by the Board as a pharmacy technician. (BPC 4115[f])
	<u>pharn</u> intran that a	pharmacy technician may, under the direct supervision and control of a nacist, prepare and administer influenza and COVID-19 vaccines via injection or nasally, prepare and administer epinephrine, perform specimen collection for tests are classified as waived under CLIA, receive prescription transfers, and accept cation on prescriptions under the following conditions: (BPC 4115[b][1])

	10.12.1. The pharmacy has scheduled another pharmacy technician to assist the pharmacist by performing the tasks as defined in BPC 4115(a), under the direct supervision of the pharmacist; (BPC 4115[b][1][A])
	□ 10.12.2. The pharmacy technician is certified and maintains the certification, by a pharmacy technician certifying organization offering a pharmacy technician certification program accredited by the National Commission for Certifying Agencies that is approved by the board; (BPC 4115[b][1][B], BPC 4202[a][4])
	□ 10.12.3. The pharmacy technician has completed at least six hours of practical training approved by the Accreditation Council for Pharmacy Education and includes hands-on injection technique, the recognition and treatment of emergency reactions to vaccines, and an assessment of the pharmacy technician's injection technique; (BPC 4115[b][1][C]; and
	☐ 10.12.4. The pharmacy technician is certified in basic life support. (BPC 4115[b][1][D])
CORRE	CTIVE ACTION OR ACTION PLAN:
	
11. Duti	es of Non-Licensed Personnel
Yes No N/A	11.1. A non-licensed person (clerk/typist) is permitted to type a prescription label or otherwise enter prescription information into a computer record system, and at the direction of a pharmacist, may request and receive refill authorization. (BPC 4007, CCR 1793.3)
	11.2. The number of non-licensed personnel supervised by each pharmacist does not interfere with the effective performance of the pharmacist's responsibilities under the Pharmacy Law. (CCR 1793.3[b])
CORRE	CTIVE ACTION OR ACTION PLAN:

PHARMACY PRACTICE

12. Pharmaceutical Service Requirements

Yes No N/A	12.1. The pharmacy complies with the requirements of 22 CCR 70263, addressing the following areas in written policies and procedures:				
	 12.1.1. Basic information concerning investigational drugs and adverse drug reactions; 12.1.2. Repackaging and compounding records; 12.1.3. Physician orders; 12.1.4. Wards, nursing stations and night stock medications; 12.1.5. Drugs brought into the facility by patients for storage or use; 12.1.6. Bedside medications; 12.1.7. Emergency drug supply; 12.1.8. Pass medications; 12.1.9. Inspection of ward stock, nursing stations and night lockers no less frequently than every 30-days\\Outdated drugs; 12.1.10. Routine distribution of inpatient medications; 12.1.11. Preparation, labeling and distribution of IV admixtures and cytotoxic agents; 12.1.12. Handling of medication when pharmacist not on duty; and 12.1.13. Use of electronic image and data order transmissions. 				
Yes No N/A □□□	12.2. The pharmacy complies with the requirements of 22 CCR 70263, addressing the following areas:				
	☐ 12.2.1. Destruction of controlled substances; and				
	$\ \square$ 12.2.2. Development and maintenance of the hospital's formulary. (22 CCR 70263)				
CORREC	CTIVE ACTION OR ACTION PLAN:				
	cation/Chart Order				
Yes No N/A	13.1. The pharmacy receives the original, the electronic transmission, or a copy of the medication order. Faxed copies, tele-autograph copies, or transmissions between computers are permissible. (BPC 688, 4019, 4040, CCR 1717.4)				
	13.2. The chart or medical record of the patient contains all of the information required by BPC 4040 and the chart order is signed by the practitioner authorized by law to prescribe drugs if present or, if not present, within a specified time frame not exceeding 48 hours. (BPC 688, 4019, 4040, 22 CCR 70263[g])				

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Yes No N/A	13.4. The pharmacy furnishes dangerous drugs or dangerous devices pursuant to preprinted or electronic standing orders, order sets and protocols established under policies and procedures. (BPC 4119.7)			
CORREC	CTIVE ACTION OR ACTION PLAN:			
14. Labe	ling and Distribution			
	14.1. Unit dose medication are properly labeled and include the information as required by BPC 4076, or the information is otherwise readily available at the time of drug administration. (BPC 4076[b])			
	14.2. The pharmacist is responsible for the proper labeling, storage and distribution of investigational drugs pursuant to the written order of the investigator. (22 Cal. Code of Regs 70263[o]).			
	14.3. This pharmacy furnishes dangerous drugs in compliance with BPC 4126.5 only to a patient pursuant to a prescription, a wholesaler from whom the dangerous drugs were purchased, a manufacturer from whom the drugs were purchased, a licensed wholesaler acting as a reverse distributor, another pharmacy to alleviate a temporary shortage with a quantity sufficient to alleviate the temporary shortage, a health care provider authorized to receive drugs, to another pharmacy of common ownership, or to a patient or to another pharmacy pursuant to a prescription. (BPC 4126.5[a])			
CORREC	CTIVE ACTION OR ACTION PLAN:			
	 			
15. Dura	tion of Drug Therapy			
Yes No N/A	15.1. The hospital has policies limiting the duration of drug therapy in the absence of the prescriber's specific indication of duration of drug therapy or under other circumstances recommended by the pharmacy and therapeutics committee or its equivalent and approved by the executive committee of the medical staff. Limitations are established for classes of drugs and/or individual drug entities. (22 CCR 70263[j])			
CORREC	CTIVE ACTION OR ACTION PLAN:			
16. Conf	identiality of Chart Orders, Prescriptions and Patient Medical Information			
Yes No N/A	16.1. Patient information is maintained to safeguard confidentiality. (Civil Code 56 et seq.)			
	16.1. Patient information is maintained to saleguard confidentiality. (Civil Code 36 et seq.) 16.2. Patient medical information, all prescriptions (chart orders, patient discharge and employee prescriptions) are confidential and are not disclosed unless authorized by law. (BPC 4040, CCR 1764, Civil Code 56 et seq.)			

Yes No N/A		estruction or disposal of patient records preserves the confidentiality of the mation contained therein. (Civil Code 56.101)	
	disc	ne pharmacy ensures electronically transmitted prescriptions (chart orders, harge patient or employee prescriptions) are received, maintained and transmitted secure and confidential manner. (BPC 688, CCR 1717.4)	
	phai	ecords regarding dangerous drugs and dangerous devices stored off-site (only for macies who have obtained a waiver from the Board of Pharmacy to store records ite) are secure and retrievable within two business days. (BPC 4105, CCR 1707)	
	Date	Waiver Approved Waiver Number	
		ress of offsite storage location:	
¥es No N/A	16.6. Records for non-controlled substances are maintained on the licensed premises for at least one year from the date of dispensing. Records for controlled substances are maintained on the licensed premises for at least two years from the date of dispensing. (BPC 4105, CCR 1707)		
CORREC	TIVE AC	CTION OR ACTION PLAN:	
17. Quali	ty Assu	rance and Medication Errors	
Yes No N/A	erro	narmacy has established quality assurance program that documents medication rs attributable, in whole or in part, to the pharmacy or its personnel. (BPC 4125, 8 1711)	
	17.2. Pharmacy quality assurance policies and procedures are maintained in the pharmacy and are immediately retrievable. (CCR 1711[c])		
	17.3. When a medication error has occurred (drug was administered to or by the patient, or resulted in a clinically significant delay in therapy) the pharmacist communicates with the patient or patient's agent that a medication error has occurred and the steps required to avoid injury or mitigate the error. (CCR 1711[c][2][A], 1711 [c][3])		
	17.4. When a medication error has occurred (drug was administered to or by the patient, or resulted in a clinically significant delay in therapy) the pharmacist communicates to the prescriber that a medication error has occurred. (CCR 1711[c][2][B], 1711 [c][3])		
	17.5. Investigation of pharmacy medication errors is initiated within two business days from the date the medication error is discovered. (CCR 1711[d])		
		ne record for quality assurance review for a medication error contains: R 1711[e]);	
		17.6.1. Date, location, and participants in the quality assurance review;	
		17.6.2. Pertinent data and other information related to the medication error(s) reviewed;	
		17.6.3. Findings and determinations;	

		17.6.4. Recommended changes to pharmacy policy, procedure, systems or processes, if any.
Yes No N/A	and	he record of the quality assurance review is immediately retrievable in the pharmacy is maintained in the pharmacy for at least one year from the date it was created. R 1711[f])
	the	harmacists are not deviating from the requirements of a prescription except upon prior consent of the prescriber, and selection of the drug product is in accordance BPC 4073 (generic substitution). (CCR 1716)
	<u>unli</u>	ne PIC is reporting the quality assurance review reports for medication errors for all censed ADDS to the Board at the time of annual renewal of the hospital pharmacy use. (CCR 1711[f])
CORRE	CTIVE A	CTION OR ACTION PLAN:
18. Rec	ord Keep	ing Requirements
Yes No N/A		ll completed pharmacy self-assessments are on file in the pharmacy and are
	mai	ntained for three years. (CCR 1715)
	for a or p of a	Il drug acquisition and disposition records (complete accountability) are maintained at least three years. Any record maintained electronically, the pharmacist-in-charge harmacist on duty is able to produce a hardcopy and electronic copy of all records equisition and disposition or other drug or dispensing-related records maintained etronically. These records include:
		18.2.1. Prescription records (BPC 4081[a])
		18.2.2. Purchase Invoices and sales records for all prescription drugs (BPC 4081)
		18.2.3. Biennial controlled substances inventory (21 CFR 1304.11)
		18.2.4. U.S. Official Order Forms (DEA Form-222) (21 CFR 1305.13, 21 CFR 1305.22)
		18.2.5. Power of Attorney for completion of DEA 222 forms (21 CFR 1305.05)
		18.2.6. Theft and Loss Reports (DEA Form 106) (21 CFR 1301.74[c])
		18.2.7. Record documenting return of drugs to wholesaler or manufacturer (BPC 4081)
		18.2.8. Record documenting transfers or sales to other pharmacies, and prescribers, and reverse distributors. (BPC 4059, 4081, 4105, 4332, CCR 1718)
		18.2.9. Centrally stored unused medications donated to a drug repository and distribution program. (HSC 150200, 150202[a][1], 150204[k], BPC 4105[c]).
	<u> </u>	18.2.10. Records documenting, to the extent possible, the kind and amounts of COVID-19 oral therapeutics furnished following a positive test for SARS-CoV-2, as

well as information regarding any testing services provided, in the patient's record in the record system maintained by the pharmacy. (BPC 4052.04[d])

Yes No N/A	18.3. Transfers or sales to other pharmacies and prescribers do not exceed five percent of the pharmacy's total annual purchases of dangerous drugs or devices. If more than five percent, registration with the board as a wholesaler has been obtained. (21 CFR 1307.11, Drug Supply Chain Security Act (DSCSA), BPC 4160)
	18.4. If sales or distributions of controlled substances to other hospitals, pharmacies, or prescribers exceed five percent of the total number of controlled substances dosage units (that are furnished to the inpatients or dispensed on prescriptions to discharge patients or employees) per calendar year, the following have been obtained: a separate DEA distributor registration and a wholesaler's permit from the board. (21 CFR 1307.11, DSCSA, BPC 4160)
	18.5. A controlled substances inventory is completed biennially (every two years).
	Date completed: (21 CFR 1304.11)
Yes No N/A	18.6. All completed controlled substances inventories are available for inspection for three years. (CCR 1718)
	 Separate Schedule II records are maintained. This includes triplicate prescriptions, invoices, US official order forms and inventory records. (21 CFR 1304.04)
	18.8. Inventories and records for Schedule III-V controlled substances are filed separately or maintained in a readily retrievable manner that distinguishes them from other ordinary business records. (21 CFR 1304.04)
	18.9. DEA Forms 222 are properly executed. (21 CFR 1305.12)
	18.10. When the pharmacy distributes Schedule II controlled substances to other DEA registrants, Copy 2 of the DEA Form 222, properly completed, are submitted at the end of each month to the DEA Regional Office. (21 CFR 1305.13)
	18.11. Any controlled substances drug loss is reported upon within one business day of discovery to the DEA and within 30 days after the date of discovery to the Board of Pharmacy of any loss of controlled substances in one of the following categories that causes the aggregate amount of unreported losses discovered in that category, on or after the same day of the previous year, to equal or exceed: within 30 days. (21 CFR 1301.74[c], CCR 1715.6)
	18.11.1. Tablets, capsules, or other oral medication, 99 dosage units
	18.11.2. Single-dose injectable medications, lozenges, film, such as oral, buccal and sublingual, suppositories, or patches, 10 dosages units.
	18.11.3. Injectable multi-dose medications, medications administered by continuous infusion, or any other multi-dose unit not described in 18.11.1, two or more multi-dose vials, infusion bags or other containers.
Yes No N/A	18.12. Records stored off-site (only for pharmacies who have obtained a waiver from the Board of Pharmacy to store records off-site) are secure and retrievable within two business days. Records for non-controlled substances are maintained on the licensed

the pharmacy comply with the requirement for a pharmacist to initial or sign a prescription record or prescription label by recording the identity of the reviewing pharmacist in a computer system by a secure means. This computer does not pe the record to be altered after made and the record of the pharmacist's identity made the computer system is immediately retrievable in the pharmacy. (CCR 1712, 171 CORRECTIVE ACTION OR ACTION PLAN: 19. Inventory Reconciliation Report of Controlled Substances		İ	premises for at least one year from the date of dispensing. Controlled substances are maintained on the licensed premises for at least two years from the date of dispensing. (CCR 1707)
19. Inventory Reconciliation Report of Controlled Substances 19.1. The pharmacy performs periodic inventory and inventory reconciliation functions detect and prevent the loss of controlled substances. (CCR 1715.65[a]) 19.2. The pharmacist-in-charge of the pharmacy reviews all inventory and inventory reconciliation reports taken, and establishes and maintains secure methods to prever losses of controlled drugs. Written policies and procedures are developed for perform the inventory reconciliation reports required by pharmacy law. (CCR 1715.65[b]) 19.3. A pharmacy compiles an inventory reconciliation report of all federal Schedule I controlled substances at least every three months. This compilation shall require: (CC 1715.65[c]) 19.3.1. A physical count, not an estimate, of all quantities of federal Schedule II controlled substances. The biennial inventory of controlled substances required federal law may serve as one of the mandated inventories under this section in tyear where the federal biennial inventory is performed, provided the biennial inventory as taken no more than three months from the last inventory required by this se (CCR 1715.65[c][1]) 19.3.2. A review of all acquisitions and dispositions of federal Schedule II controsubstances since the last inventory reconciliation report; (CCR 1715.65[c][2]) 19.3.3. A comparison of the two above-mentioned items to determine if there are variances; (CCR 1715.65[c][3]) 19.3.4. All records used to compile each inventory reconciliation report shall be maintained in the pharmacy or clinic for at least three years in a readily retrievab form; and (CCR 1715.65[c][4]) 19.3.5. Possible causes of overages shall be identified in writing and incorporate the inventory reconciliation report. (CCR 1715.65[c][5]) 19.3.6. In addition to Schedule II controlled substance, the pharmacy is performiny enterory reconciliation of alprazolam 1mg/unit, alprazolam 2mg/unit. tramadol 50mg/unit, and promethazine with codeine 5.25mg promethazine/10mg codeine		1 	
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		Ω	19.3.6. In addition to Schedule II controlled substance, the pharmacy is performing an inventory reconciliation of alprazolam 1mg/unit, alprazolam 2mg/unit, tramadol 50mg/unit, and promethazine with codeine 6.25mg promethazine/10mg codeine per 5ml of product at least every 12 months. (CCR 1715.65[a][2])

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	☐ 19.3.7. An inventory reconciliation report must be prepared for any identified controlled substances lost no later than three months after discovery of the reportable loss. (CCR 1715.65)
	☐ 19.3.8. Inventory activities for all other controlled substances must be performed at least once every two years from the performance of the last inventory activities. (CCR 1715.65[a][3][B])
	□ 19.3.9. The inventory reconciliation report may use a digital or electronic signature or biometric identifier in lieu of a physical signature if, in addition, the individual physically signs a printed statement confirming the accuracy of the inventory or report. The signature shall be dated, and the signed and dated statement shall be retained on file. (CCR 1715.65[e][1])
	□ 19.3.10. Inpatient hospital pharmacy, the inventory reconciliation for all federal Schedule II-controlled substances, and alprazolam 1mg/unit, alprazolam 2mg/unit, tramadol 50mg/unit, and promethazine with codeine (6.25mg promethazine/10mg codeine/5ml) must be performed on a quarterly basis. The report or reports shall include controlled substances stored within the pharmacy, within each pharmacy satellite location, and within each drug storage area in the hospital under the pharmacy's control. (CCR 1715.65[g])
Yes No N/A	19.4 The pharmacy reports in writing identified losses and known causes to the board within 30 days of discovery unless the cause of the loss is theft, diversion, or self-use in which case the report shall be made within 14 days of discovery. If the pharmacy is unable to identify the cause of the loss, further investigation is undertaken to identify the cause and actions necessary to prevent additional losses of controlled substances. (BPC 4104, CCR 1715.65[d], CCR 1715.6)
Yes Ne N/A □□□□	19.5 The inventory reconciliation report is dated and signed by the individual(s) performing the inventory, and countersigned by the pharmacist-in-charge and be readily retrievable in the pharmacy for three years. A countersignature is not required if the pharmacist-in-charge personally completed the inventory reconciliation report. (CCR 1715.65 [e])
	19.65 A new pharmacist-in-charge of the pharmacy completes an inventory reconciliation report as identified in CCR 1715.65 (c) within 30 days of becoming pharmacist-in-charge. When possible, the outgoing pharmacist-in-charge also completes an inventory reconciliation report as required in CCR 1715.65 (c). (CCR 1715.65 [f])
	19.7 A separate quarterly inventory reconciliation report shall be required for federal Schedule II controlled substances stored within the pharmacy and for each pharmacy satellite location. (CCR 1715.65 [g])
	19.8 The pharmacist-in-charge of an inpatient hospital pharmacy or of a pharmacy servicing onsite or offsite automated drug delivery systems shall ensure that: (CCR 1715.65[h])
	 □ 19.8.1 All controlled substances added to an automated drug delivery system are accounted for; (CCR 1715.65[h][1]) □ 19.8.2 Access to automated drug delivery systems is limited to authorized facility personnel; (CCR 1715.65[h][2])

	☐ 19.8.3 An ongoing evaluation of discrepancies or unusual access associated with controlled substances is performed; and (CCR 1715.65[h][3])		
	□ 19.8.4 Confirmed losses of controlled substances are reported to the board. (CCR 1715.65[h][4])		
Yes No N/A	19.8. If the inpatient hospital pharmacy uses an ADDS, inventory in the ADDS may be accounted for under CCR section 1715.65(c)(1) using means other than a physical count. (CCR 1715.65[h])		
CORRE	CTIVE ACTION OR ACTION PLAN:		
	r-Hours Supply of Medication		
Yes No N/A	20.1 The pharmacy has a system assuring the prescribed medications are available in the hospital 24 hours a day. (22 CCR 70263[e])		
	20.2. The pharmacy maintains a record of the drugs taken from the after-hours supply of medications and the pharmacist is notified of such use. The record includes the name and strength of the drug, the amount taken, the date and time, the name of the patient to whom the drug was administered and the signature of the registered nurse. (22 CCR 70263[n])		
CORRE	CTIVE ACTION OR ACTION PLAN:		
21. Dru	g Supplies for Use in Medical Emergencies		
Yes No N/A	21.1. A supply of drugs for use in medical emergencies only is immediately available at each nursing unit or service area as required. (22 CCR 70263[f])		
	21.2. Written policies and procedures have been developed that establish the contents of the supply, procedures for use, restocking and sealing of the emergency drug supply. (22 CCR 70263[f][1], BPC 4115, 4119.6))		
	21.3. The emergency drug supply is stored in a clearly marked portable container which is sealed by the pharmacist in such a manner that a seal must be broken to gain access to the drugs. The contents of the container are listed on the outside cover and include the earliest expiration date of any drugs within. (22 CCR 70263[f][2])		
	21.4. The pharmacist is responsible for inspection of the drug supply at periodic intervals (no less frequently than every 30 days) specified in the writ-ten policies. Records of the inspection are kept for at least three years. The inspection can be done by a pharmacy technician or pharmacy intern as defined in the pharmacy's written inspection policies and procedures. (22 CCR 70263[f][3], BPC 4115[ji][3], 4119.7[c])		
CORRE	CTIVE ACTION OR ACTION PLAN:		

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22. Sche	dule II-	/ Controlled Substances Floor Stock Distribution Records		
Yes No N/A	22.1. Records for the distribution of Schedule II-V controlled substances floor stock are open to inspection by authorized officers of the law and are preserved for at least three years from the date of making. (BPC 4081)			
CORREC	CTIVE A	CTION OR ACTION PLAN:		
	rgency I	Room Dispensing		
Yes No N/A		prescriber may dispense a dangerous drug, including a controlled substance, to an ergency room patient if all of the following apply: (BPC 4068[a])		
		23.1.1. The hospital pharmacy is closed and there is no pharmacist available in the hospital;		
		23.1.2. The dangerous drug is acquired by the hospital pharmacy;		
		23.1.3. The dispensing information is recorded and provided to the pharmacy when the pharmacy reopens;		
		23.1.4. The hospital pharmacy retains the dispensing information and, if the drug is a schedule II, III, IV or IV controlled substance, transmits the dispensing data to the Department of Justice within one working day from the date the controlled substance is released to the patient. (HSC 11165[d])		
		23.1.5. The prescriber determines that it is in the best interest of the patient that a particular drug regimen be immediately commenced or continued, and the prescriber reasonably believes that a pharmacy located outside the hospital is not available and accessible at the time of dispensing to the patient; and		
		23.1.6. The quantity of drugs dispensed to any patient pursuant to this section are limited to that amount necessary to maintain uninterrupted therapy during the period when pharmacy services outside the hospital are not readily available or accessible, but shall not exceed a 72-hour supply;		
	Ω	23.17. If an ADDS is located in the emergency room and is used for dispensing to patients upon discharge, the ADDS is licensed with the Board. (BPC 4427.2[i]).		
Yes No N/A	23.2. The prescription label contains all the required information and is formatted in accordance with CCR 1707.5 including Patient Centered Labels in at least 12-point sans serif typeface for the four required items in the required order. (BPC 4076, CCR 1707.5)			
		he prescriber shall be responsible for any error or omission related to the drugs bensed. (BPC 4068[b])		

<u>Yes No N/A</u> □□□□	23.4. The brand name or generic name and manufacturer of the prescription drug is accurately identified on the label and prescription record. (BPC 4076, CCR 1717)
	23.5. Controlled substances are dispensed in prescription containers bearing a federal warning label prohibiting transfer of the drugs. (21 CFR 290.5)
	23.6. Prescriptions are dispensed in new, senior-adult ease-of-opening tested, and child-resistant containers or in a noncomplying package, only pursuant to the prescription or when requested by the purchaser. (15 USC 1473[b], 16 CFR 1700.15-, CCR 1717)
	23.7. Patient package inserts are dispensed with all estrogen medications (21 CFR 310.515)
	23.8. The pharmacy provides patients with required Black Box Warning Information. (21 CFR 201.57[c])
	23.9. Medication guides are provided on required medications. (21 CFR Part 208)
	23.10. Whenever an opioid prescription drug is dispensed to a patient for outpatient use, the pharmacy prominently displays on the label or container or by a flag or other notification to the container, a notice that states, "Caution: Opioid. Risk of overdose and addiction." (BPC 4076.7).
	23.11. A pharmacist may dispense a drug prescribed pursuant to HSC Section 120582 and label the drug without the name of an individual for whom the drug is intended if the prescription includes the words "expedited partner therapy" or the letters "EPT" and shall provide written notification that describes the right of an individual who received EPT to consult with a pharmacist about the medication dispensed and possible drug interactions. (BPC 4076[f], [h])
	23.12. If emergency department patient dispensing is done via AUDS, the AUDS is licensed by the Board. (BPC 4427.2[i])
	23.13. A practitioner specified in Section 11150 may dispense a controlled substance
	classified in Schedule II, which may be from a hospital pharmacy inventory, directly to an ultimate user in either of the following circumstances: (HSC 11158)
	23.13.1 In an amount not to exceed a 72-hour supply for the patient in accordance with directions for use given by the dispensing practitioner only when the patient is not expected to require any additional amount of the controlled substance beyond the 72 hours; (HSC 11158[b][1])
	23.13.2 For the purpose of initiating maintenance treatment or detoxification
	treatment, or both, for a person with an opioid use disorder. Not more than a three-day supply of such medication may be dispensed to the person at one time while
	arrangements are being made for referral for treatment. Such emergency treatment
	may not be renewed or extended. (HSC 11158[b][2])
CORREC	CTIVE ACTION OR ACTION PLAN:

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24. Discharge Medication/Consultation Services

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Yes No N/A	24.1. Patients receive information regarding each medication given at the time of discharge. The information includes the use and storage of each medication, the precautions and relevant warnings and the importance of compliance with directions. A written policy has been developed in collaboration with a physician and surgeon, a pharmacist, and a registered nurse and approved by the medical staff that ensures that each patient receives the medication consultation. (BPC 4074, CCR 1707.2)
	24.2. Prescriptions are transmitted to another pharmacy as required by law. (BPC 4072, CCR 1717[c], [f], 1717.4)
	24.3. The prescription label contains all the required information and is formatted in accordance with CCR 1707.5 including Patient Centered Labels in at least 12-point sans serif typeface for the four required items in the required order. (BPC 4076, CCR 1707.5)
Yes No N/A	24.4. The pharmacist includes a written label on the drug container indicating that the drug may impair a person's ability to operate a vehicle or vessel. The label may be printed or an auxiliary label affixed to the prescription container. (BPC 4074 [a], [b], CCR 1744[a][1]-[7])
	24.5 The pharmacist includes a written label on the drug container to alert the patient about possible potentiating effects when taken in combination with alcohol. The label may be printed on an auxiliary label affixed to the prescription container (BPC 4074[a], CCR 1744[b][1]-[6]).
	24.6. The trade name or generic name and manufacturer of the prescription drug is accurately identified in the prescription record. (CCR 1717)
	24.7. Generic substitution for discharge medications is communicated to the patient, and the name of the dispensed drug product is indicated on the prescription label. (BPC 4073)
	24.8. If the prescription is filled by a pharmacy technician, the pharmacist's initials are on the prescription label to document the pharmacist's verification of the product or can be satisfied by recording the identity of the reviewing pharmacist in a computer system by a secure means and is immediately retrievable in the pharmacy. (CCR 1712, 1793.7)
	24.9. Controlled substances are dispensed in prescription containers bearing a federal warning label prohibiting transfer of the drugs. (21 CFR 290.5)
	24.10. Prescriptions are dispensed in a new and child-resistant container, or senior-adult ease-of-opening tested container, or in a non-complying package only pursuant to the prescriber or when requested by the purchaser. (15 USC 1473, 16 CFR 1700.15, CCR 1717)
	24.11. Patient package inserts are dispensed with all estrogen medications. (21 CFR 310.515)
Yes No N/A	

	24.12. The pharmacy provides patients with required Black Box Warning. (21 CFR 201.57[c])
	24.13. Medication guides are provided on required medications. (21 CFR Part 208)
	24.14. Whenever an opioid prescription drug is dispensed to patient for outpatient use, the pharmacy prominently displays on the label or container or by a flag or other notification to the container, a notice that states, "Caution: Opioid. Risk of overdose and addiction." (BPC 4076.7).
	24.15. Effective January 1, 2022, t The pharmacy has the capability to receive electronic data transmission prescriptions on behalf of patients. (BPC 688)
	24.16. The hospital pharmacy retains the dispensing information and, if the drug is a schedule II, III, IV or V controlled substance, transmits the dispensing data to the Department of Justice within one working day after the date the controlled substance is released to the patient or patients representative. (HSC 11165[d])
	24.17. Drugs dispensed pursuant to a veterinary prescription shall include, as part of the consultation, the option for a representative of an animal patient to also receive drug documentation specifically designed for veterinary drugs. (BPC 4069)
CORREC	CTIVE ACTION OR ACTION PLAN:
	ral Filling of Patient Cassettes For Other Hospital Pharmacies
Yes No N/A	25.1. Pharmacy processes orders for the filling of patient cassettes for another hospital or Pharmacy within this state receives filled medication orders or patient cassettes from another hospital. (CCR 1710[b])
	If the answer is "yes," name of hospital:
	25.2. Pharmacy receives filled medication containers or cassettes from another pharmacy. (CCR 1710[b])
	If the answer is "yes," name of supplying pharmacy:
	If the answer to this and the previous question is "no" or "not applicable" go to Section 26.
	25.3. Prescription information is electronically transferred between the two pharmacies. (CCR 1710[b][6])
	25.4. Pharmacy has a contract with the ordering hospital pharmacy or has the same owner (CCR 1710[b][1])
	25.5. Filled cassettes are delivered directly to the ordering hospital pharmacy. (CCR 1710[b][2])
	25.6. Each cassette or container meets the requirements of Business and Professions Code section 4076. (BPC 4076[b], [c], [d], CCR 1710[b][3])
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	includ	Complete and accurate records are maintained of each cassette fill transaction, uding the name of the pharmacist checking the cassettes at each pharmacy. CR 1710[b][5])		
	ralized Ho	ospital I	Packaging Pharmacy	
Yes No N/A	hosp	ital phar		ackaging, the pharmacy in addition to the Centralized Hospital Packaging specialty
Lic	ense Nur	mber:		
	funct and c	ions, for one or m	administration only to inpatien	performing the following specialized ts within its own general acute care hospital ils under common ownership and located
	Hosp	itals to v	which central packaged unit do	se medications are provided:
	□ 26	6.2.1.		Distance (miles):
	□ 26	6.2.2.		Distance (miles):
	□ 26	6.2.3. _.		Distance (miles):
	□ 26	6.2.4.		Distance (miles):
				single administration to inpatients from bulk barcoded pursuant to BPC 4128.4.
			•	it dose drugs for administration to inpatients arcoded pursuant to BPC 4128.4.
			repares compounded unit dose dose package is barcoded pur	drugs for administration to inpatients, if suant to BPC 4128.4.
Yes No N/A		ient-spe	• • •	d quantities of unit dose drugs in advance or ecessary to ensure continuity of care. (BPC
	26.4. Any unit dose medications produced by a centralized ho are barcoded to be machine readable at the inpatient's be medication administrative software. (BPC 4128.4)		inpatient's bedside using barcode	
		practitio the right	ners to ensure that before a m	istration software permits health care edication is administered to an inpatient, it is ent, in the right dose, and via the right route
		reading		nedication satisfies the above criteria by and comparing the information retrieved to patient. [BPC 4128(b)]

	26.54. Any label for each unit dose medication produced by a centralized hospital packaging pharmacy displays a human-readable label that contains the following: (BPC 4128.5[a])				
		 26.5.1. The date the medication was prepared. 26.5.2. The beyond-use date 26.5.3. The established name of the drug. 26.5.4. The quantity of each active ingredient. 26.5.5. The lot number or control number assigned by the centralized hospital packaging pharmacy. 26.5.6. Special storage or handling requirements. 26.5.7. The name of the centralized hospital packaging pharmacy. 			
Yes No N/A		he pharmacist is able to retrieve all of the following information using the lot number ontrol number: (BPC 4128.5[b])			
		26.6.1. The components used in the drug product.26.6.2. The expiration date of each of the drug's components.26.6.3. The National Drug Code Directory number.			
	pha unit	he centralized hospital packaging pharmacy and the pharmacists working in the rmacy are responsible for the integrity, potency, quality, and labeled strength of any dose drug product prepared by the centralized hospital packaging pharmacy. C 4128.7)			
CORREC	CTIVE A	CTION OR ACTION PLAN:			
	cies and	Procedures			
Yes No N/A □□□□	27.1. T	here are written policies and procedures in place for:			
		27.1.1. Oral consultation for discharge medication to an inpatient of a health care facility licensed pursuant to HSC 1250. The assurance that each patient received information regarding each medication given at the time of discharge. (BPC 4074[e], CCR 1707.2[b][2])			
		27.1.2. Action to be taken to protect the public when a licensed individual employed by or with the pharmacy is known to be chemically, mentally, or physically impaired to the extent that it effects their ability to practice the profession or occupation authorized by their license. (BPC 4104[a])			
		27.1.3. Action to be taken to protect the public when a licensed individual employed by or with the pharmacy is known to have engaged in the theft or diversion or self-use of prescription drugs belonging to the pharmacy. (BPC 4104[b])			
		27.1.4. Addressing chemical, mental, or physical impairment, as well as, theft, diversion, or self-use of dangerous drugs, among licensed individual employees by or with the pharmacy. (BPC 4104[b])			

	information as specified in BPC 4104[c][1]-[6].
	27.1.6. Operation of the pharmacy during the temporary absence of the pharmacist for breaks and meal periods including authorized duties of personnel, pharmacist's responsibilities for checking all work performed by ancillary staff, and pharmacist's responsibility for maintaining the security of the pharmacy. (CCR 1714.1[f])
	27.1.7. Assuring confidentiality of medical information if your pharmacy maintains the required dispensing information for prescriptions, other than controlled substances, in a shared common electronic file. (CCR 1717.1[e])
	27.1.8. Helping patients with limited or no English proficiency understand the information on the prescription container label in the patient's language, including the selected means to identify the patient's language and providing interpretive services in the patient's language. (CCR 1707.5)
	27.1.9. Inventory reconciliation reporting requirements. (CCR 1715.65)
	27.1.10. Pharmacy technician performing monthly checks of the drug supplies stored throughout the health care facility and reporting irregularities within 24 hours to the pharmacist-in-charge and the director or chief executive officer of the health care facility. (BPC 4115[ji][3])
	27.1.11. Intern pharmacist, under the direct supervision and control of a pharmacist, may inspect the drugs maintained in the health care facility at least once per month. (BPC 4119.7[c])
	27.1.12. Furnishing dangerous drug or dangerous device pursuant to preprinted or electronic standing orders, order sets, and protocol, if the order is dated, timed, and authenticated in the medical record of the patient to whom the dangerous drug or dangerous device is provided. (BPC 4119.7[a])
	27.1.13. Storing and maintaining drugs in accordance with national standards regarding storage areas, refrigerator or freezer temperature, and otherwise pursuant to the manufacturer's guidelines. (BPC 4119.7[b], 22 CCR 70263 [c][1], [q][6])
	27.1.14. Establishing the supply contents, procedure for use, restocking and sealing of emergency drug supply. (CCR 70263[f][1])
	27.1.15. If applicable, dispensing, storage and records of use if bedside medications are allowed. No controlled substances shall be left at bedside. (22 CCR 70263[/])
	27.1.16. The use of investigational drugs. Basic information concerning the dosage form, route of administration, strength, actions, uses, side effects, adverse effects, interaction and symptoms of toxicity shall be available in the pharmacy and the nursing station. The pharmacist is responsible for the proper labeling, storage and distribution of such drug pursuant to the investigator's written orders. (22 CCR 70263[o]).
CORRECTIVE A	CTION OR ACTION PLAN:
	
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27.1.5. Reporting to the board within 14 days of the receipt or development of

28. Com	pounding		
Yes No N/A	Prior to allowing any drug product to be compounded in a pharmacy, the pharmacist-in-charge must complete the "Compounding Self-Assessment" as required by CCR 1735.2. (CCR 1735.2)		
29. Auto	omated Drug Delivery Systems		
Yes No N/A	29.1. The hospital pharmacy operates automated drug delivery systems that are automated unit dose systems (AUDS) for doses administered at the facility and approved services listed on the hospital's license and the ADDS is/are exempt from licensure with the board. The AUDS must comply with all other requirements for an ADDS in Article 25. (BPC 4427.2[i])		
	29.2. The hospital pharmacy operates automated drug delivery systems that are automated patient delivery dispensing systems (APDS) for doses dispensed to patients at the facility and approved services listed on the hospital's license and the ADDS is/are licensed with the board. (BPC 4427.2[a])		
	29.3. If the pharmacy operates an automated drug delivery system, the pharmacist-in-charge has completed the self-assessment for automated drug delivery systems pursuant to CCR 1715. The pharmacy shall comply with all recording keeping and quality assurance requirements and maintain those records within the licensed pharmacy holding the ADDS license and separate from other pharmacy records. (BPC 4427.7)		
	CTIVE ACTION OR ACTION PLAN:		
Yes No N/A	30.1. Does the pharmacy participate in a Prescription Drug Take-Back Program and adheres to the federal, state, and local requirements governing the collection and destruction of dangerous drugs? (CCR 1776, 1776.1)		
	If yes, check off below the type of prescription drug take-back program the pharmacy offers and complete the sections that apply to the type of program(s):		
	 □ Mail back envelopes or package service. (CCR 1776.2) □ Collection receptacles in the pharmacy. (CCR 1776.3) □ Drug take-back services in the Skilled Nursing Facilities. (CCR 1776.4, HSC 1250[c]) 		
	30.2. Only prescription drugs that have been dispensed by any pharmacy or practitioner to a consumer are eligible for collection as part of drug take-back services maintained by the pharmacy. (CCR 1776.1[f])		
	30.3. Dangerous drugs that have not been dispensed to consumers for use (such as outdated drug stock, drug samples provided to medical practitioners or medical waste) are not collected as part of the pharmacy's drug take-back service. (CCR 1776.1[f])		
Yes No N/A			

Pharma Yes No N/A	cies with Collection Receptacles in the Pharmacy/Hospital (CCR 1776.1, 1776.3)
CORRE	the pharmacy does not remove, count, sort or individually handle any prescription drugs from the consumer. (CCR 1776.1[d], [g]) CTIVE ACTION OR ACTION PLAN:
	Expiration Date: 30.11. Once drugs are deposited into a mail back envelope or package by the consumer,
	DEA Collector Registration Number:
	CFR 1317.40):
	If the answer is no and the pharmacy is registered with DEA as a collector with an onsite method of destruction that complies with DEA requirements, list the following (21
	30.10. The pharmacy does not accept any mail back packages or envelopes containing drugs unless the pharmacy is registered as a collector and has an onsite method of destruction that complies with DEA requirements. The consumer is directed to mail the envelopes or packages. (CCR 1776.2[e])
Yes No N/A	30.9. The preaddressed envelope and package contains a unique identification number for each envelope and package, and the instructions for users indicates the process to mail back the drugs. (CCR 1776.2[d])
Vac No N/A	30.8. The preaddressed envelopes and packages are water and spill proof, tamper evident tear resistant and sealable. The exterior is nondescript and has no markings indicating the envelope or package contains prescription drugs. Postage is prepaid on each envelope or package. (CCR 1776.2[c])
	30.7. All drug take-back envelopes and packages made available to the public are preaddressed to a location registered with the DEA as a collector. (CCR 1776.2[b])
Yes No N/A	30.6. The pharmacy provides prescription drug take-back preaddressed mailing envelopes or packages to consumers to return prescription drugs to an authorized destruction location. (CCR 1776.2[a])
	cies Offering Mail Back Envelopes or Package Services (CCR 1776.1, 1776.2)
CORRE	CTIVE ACTION OR ACTION PLAN:
	30.5. Quarantined, recalled or outdated prescription drugs from the pharmacy stock are not disposed as part of the pharmacy's drug take-back services. (CCR 1776.1[g][3])
	its drug take-back services. (CCR 1776.1[g][2])
	30.4. The pharmacy does not accept or possess prescription drugs from skilled nursing facilities, residential care homes, health care practitioners or any other entity as part of

	30.12. The pharmacy is registered with DEA as a collector for purposes of maintaining a prescription drug take-back collection receptacle. (21 CFR 1317.30, 1317.40, CCR 1776)		
	30.13. The pharmacy notified the board in writing within 30 days of establishing the collection program. (CCR 1776.1[i])		
	Date the board was notified:		
	30.14. When the pharmacy renewed its pharmacy license, the pharmacy identified and disclosed to the board all the locations where its collection receptacles are located. (CCR 1776.1[i][2])		
	30.15. The pharmacy is aware, any tampering with a collection receptacle or theft of deposited drugs and any tampering, damage or theft of the removed liner are reported to the board in writing within 14 days. (CCR 1776.1[i][3][4])		
	List the dates the board was notified of any tampering or theft from the collection receptacle and/or tampering, damage or theft of the removed liner:		
	Date reported:		
	30.16. The pharmacy is not on probation with the board. (CCR 1776.1[I])		
	If answered NO, meaning the pharmacy is on probation, the pharmacy cannot maintain a drug take back collection receptacle and must cease and notify the board in writing within 30 days and notify the DEA within 30 days.		
Yes No N/A	30.17. Once drugs are deposited into a collection receptacle by the consumer, the pharmacy does not remove, count, sort or individually handle any prescription drugs from the consumer. (CCR 1776.1[d], 1776.3[e])		
	30.18. The collection receptacle is substantially constructed with a permanent outer container, removable inner liner, and is locked at all times to prevent access to the inner liner. (CCR 1776.3[a], [d])		
	30.19. The collection receptacle is securely fastened to a permanent structure so it cannot be removed and is installed in an inside location. (CCR 1776.3[b])		
	30.20. The receptacle is visible to the pharmacy and DEA registrant employees, but not located in or near emergency areas, nor behind the pharmacy's counter or is located in an area that is regularly monitored by pharmacy or DEA registrant employees and not in the proximity of any emergency or urgent care areas. When no pharmacy or DEA registrant employees are present, the collection receptacle is locked so that drugs are not deposited into the collection receptacle. (CCR 1776.3[b], [c])		
	30.21. The receptacle includes a small opening that allows deposit of drugs into the inside of the receptacle directly into the inner liner, but does not allow for an individual to reach into the receptacle's contents. When the pharmacy is closed, the collection receptacle is not accessible to the public for deposit of drugs. The pharmacy locks the deposit opening on the collection receptacle. (CCR 1776.3[d])		
Yes No N/A	30.22. The pharmacy directs consumers to directly deposit the drugs into the collection receptacle. (CCR 1776.3[e])		

	30.23. The inner liner used is made of material that is certified by the manufacturer to meet the ASTM D179 standard test for impact resistance of 165 grams (drop dart test) and the ASTM D1922standards for tear resistance of 480 grams in both parallel and perpendicular planes. (CCR1776.3[f])		
	 □ 30.23.1 The liner is waterproof, tamper evident, tear resistant, and opaque to prevent viewing or removal of any contents once the liner has been removed from the collection receptacle. (CCR 1776.3[f]) □ 30.23.2 The liner is clearly marked to display the maximum contents (for example, 		
	in gallons). (CCR 1776.3[g]) 30.23.3 The liner bears a permanent, unique identification number established by the pharmacy or pre-entered onto the liner by the liner's manufacturer or distributor. (CCR 1776.3[f][2])		
	□ 30.23.4 The liner is removable as specified pursuant to CCR 1776.3. (CCR 1776.3[f][2])		
	30.24. The receptacle allows the public to deposit prescription drugs into the receptacle for containment into the inner liner, without permitting access to or removal of prescription drugs already deposited into the collection receptacle and liner. Once the prescription drugs or any other item is placed in the collection receptacle, the prescription drug or item cannot be removed, counted, sorted, or otherwise individually handled. (CCR 1776.3[d], [e], [g])		
Yes No N/A	30.25. If the liner is not already itself rigid or already inside of a rigid container when it is removed from the collection receptacle, the liner is immediately, without interruption, placed in a rigid container for storage, handling and transport. (CCR 1776.3[h])		
	30.26. The liner is removed from a locked collection receptacle only by or under the supervision of two employees of the pharmacy. Upon removal, the liner is immediately, without interruption, sealed and the pharmacy employees record, in a log, their participation in the removal of each liner from the collection receptacle. Liners and their rigid containers are not opened, x-rayed, analyzed, or penetrated at any time by the pharmacy or pharmacy personnel. (CCR 1776.3[i])		
	30.27. Liners and their rigid containers that are filled and removed from the collection receptacle is stored in a secured, locked location in the pharmacy no longer than 14 days. (CCR 1776.3[j])		
	30.28. The pharmacy maintain records for collected unwanted drugs from consumers for three years, including the records for each liner. (CCR 1776.3[k], 1776.6[a])		
	30.29. The pharmacy seals the inner liners and their contents are shipped to a reversed distributor's registered location by common or contract carrier (such as UPS, FEDEX, USPS) or by a licensed reverse distributor pick up at the licensed pharmacy's premise. (CCR 1776.3[I])		
Yes No N/A	30.30. The collection receptacle has a signage that includes: (1) the name and phone number of the responsible pharmacy, (2) medical sharps and needles (e.g. insulin syringes) shall not to be deposited, and (3) consumers may deposit prescription drugs including Schedule II-V controlled substance. (CCR 1776.1[e], 1776.3[m])		
CORRECTIVE ACTION OR ACTION PLAN:			
	DIC.		

Onsite Pharmacies with Drug Take-Back Services in Skilled Nursing Facilities Yes No N/A 30.31. The pharmacy provides mail back envelopes or packages to skilled nursing facility employees or persons lawfully entitled to dispose of a resident decedent's property of unwanted or unused prescription drugs. (CCR 1776.4[a]) 30.32. If the pharmacy provides mail back envelopes or packages to skilled nursing facilities, the skilled nursing facility employees keeps a record noting the specific quantity of each prescription drug mailed back, the unique identification number of the mail back package and the preaddressed location to which the mail back envelope is sent. (CCR 1776.4[a]) 30.33. If the pharmacy is onsite at a skilled nursing facility, has the pharmacy established a collection receptacle in the skilled nursing facility for the collection and ultimate disposal of unwanted prescription drugs? (CCR 1776.4[b]) If no, answer N/A to the remaining questions in this section. If yes, continue answering the questions in this section. List the location(s) of the collection receptacle: Yes No N/A 30.34. The board was notified in writing within 30 days of establishing a collection receptacle. (CCR 1776.4[b][2]) 30.35. Has there been any tampering of the collection receptacle, theft of the deposited drugs and/or tampering, damage or theft of the removed liner? (CCR 1776.4[b][4], [5]) If yes, was the board notified in writing within 14 days of any tampering of the collection receptacles and/or liner? 30.36. When the pharmacy license was renewed, the pharmacy provide the list a current list of collection receptacles. (CCR 1776.4[b][6]) 30.37. The skilled nursing facility places a patient's unneeded prescription drugs into a collection receptacle within three business days after the permanent discontinuance of use of a medication by a prescriber, as a result of the resident's transfer to another facility or as a result of death. Records of such deposit is made in the patient's records, with the name and signature of the employee discarding the drugs. (CCR 1776.4[d]) Yes No N/A 30.38. The collection receptacle is located in a secured area regularly monitored by the skilled nursing facility employees, securely fastened to a permanent structure so it cannot be removed, has a small opening that allows deposit of drugs into the inside of the collection receptacle and directly into the inner liner, but does not allow for an individual to reach into the receptacle's contents, securely locked and substantially

	constructed with a permanent outer container and a removable inner liner. (CCR 1776.4[e][f][g])
	30.39. The liner is certified by the manufacturer to meet the American Society for Testing Materials(ASTM) D1709 standard test for impact resistance of 165 grams (drop dart test) and the ASTM D1922standards for tear resistance of 480 grams in both parallel and perpendicular planes, is waterproof, tamper evident, tear resistant, and opaque to prevent viewing and discourage removal of any contents once the liner is removed from the collection receptacle, marked to display the maximum contents, and bear a permanent, unique identification number. (CCR 1776.4[h])
	30.40. The receptacle allows the deposit of prescription drugs into the receptacle for containment into the inner liner, without permitting access to or removal of prescription drugs already deposited into the collection receptacle and liner. Once a prescription drug or item is placed in the collection receptacle, the prescription drug or item cannot be removed, sorted, counted, or otherwise individually handled. (CCR 1776.4[g][1])
	30.41. If the liner is not already itself rigid or already inside a rigid container when it is removed from the collection receptacle, the liner is immediately placed in a rigid container for storage, handling and transport. (CCR 1776.4[g][2])
Yes No N/A	30.42. The rigid container is either disposable, reusable, or recyclable. The rigid container is leak resistant, has sealable tight fitting covers and kept clean and in good repair. (CCR 1776.4[g][2])
	30.43. The collection receptacle contains signage with (1) the name and phone number of the pharmacy, (2) medical sharps and needles (e.g. insulin syringes) can not be deposited, and (3) consumers may deposit prescription drugs including Schedule II-V controlled substances. (CCR 1776.4[i])
	30.44. Once deposited, the prescription drugs are not counted, sorted, or otherwise individually handled. (CCR 1776.4[j])
	30.45. The installation, removal, transfer, and storage of inner liners is performed only by (1) one employee of the authorized collector pharmacy and one supervisory level employee of the long term care facility (e.g. charge nurse or supervisor) designated by the authorized collector or (2) by or under the supervision of two employees of the authorized collector pharmacy. (CCR 1776.4[k])
	30.46. Sealed inner liners placed in a container are stored at the skilled nursing facility up to three business days in a securely locked, substantially constructed cabinet or a securely locked room with controlled access until transfer to a reverse distributor for destruction. (CCR 1776.4[I])
Yes No N/A	30.47. Liners housed in a rigid container are delivered to a reverse distributor for destruction by a common or contract carrier or by a reverse distributor picked up at the skilled nursing facility. (CCR 1776.4[m])
CORREC	CTIVE ACTION OR ACTION PLAN:

Record P	Ceeping	Requirements for Board Licensees Providing Drug Take Back Services
Yes No N/A	30.48. F	Records required for drug take back services are maintained for three years. (CCR 5.6)
		The pharmacy makes and keeps the following records for each liner: (CCR 6.6[a])
		30.49.1. The date each unused liner is acquired, its unique identification number and size (e.g. 5 gallon, 10 gallon). If the liner does not already contain a unique identification number, the pharmacy assigns the unique identification number. (CCR 1776.6[a][1])
		30.49.2. The date each liner is installed in a collection receptacle, the address of the location where each liner is installed, the unique identification number and size (e.g., 5 gallon, 10 gallon), the registration number of the collector pharmacy, and the names and signatures of the two employees that witnessed each installation. (CCR 1776.6[a][2])
		30.49.3. The date each inner liner is removed and sealed, the address of the location from which each inner liner is removed, the unique identification number and size (e.g. 5 gallon, 10 gallon) of each inner liner removed, the registration number of the collector pharmacy, and the names and signatures of the two employees that witnessed the removal and sealing. (CCR 1776.6[a][3])
		30.49.4. The date each sealed inner liner is transferred to storage, the unique identification number and size (e.g., 5 gallon, 10 gallon) of each inner liner stored, and the names and signatures of the two employees that transferred each sealed inner liner to storage. (CCR 1776.6[a][4])
		30.49.5. The date each sealed inner liner is transferred for destruction, the address and registration number of the reverse distributor or distributor to whom each sealer inner liner was transferred, the unique identification number and size (e.g., 5 gallon, 10 gallon) of each liner transferred, and the names and signatures of the two employees who transferred each sealed inner liner to the reverse distributor or distributor, or the common carrier who delivered it, the company used, and any related paperwork (invoice, bill of lading). (CCR 1776.6[a][5])
CORREC	CTIVE AC	CTION OR ACTION PLAN:

PIC

PHARMACIST-IN-CHARGE CERTIFICATION:

	DDU //	
I, (please print) that I have completed the self-assessment of this Any deficiency identified herein will be corrected responses are subject to verification by the Board	by (date). I	l understand that all
perjury of the laws of the State of California that t assessment form is true and correct.		
Signature		Date
(Pharmacist-in-Charge)		
ACKNOWLEDGEMENT BY HOSPITAL ADMIN	ISTRATOR:	
I, (please print) the laws of the State of California that I have read understand that failure to correct any deficiency is identified in the Pharmacist-in-Charge Certification pharmacy's license issued by the California State	dentified in this self-asses on above could result in the	sment in the timeframe
Signature		Date
(Hospital Administrator)		

The following **Legal References** are used in the self-assessment form. Many of these references can be viewed on the Board of Pharmacy Web site at www.pharmacy.ca.gov (see *Laws and Regulations*), at the California State Law Library, or at other libraries or Internet web sites.

- Business and Professions Code (BPC), Division 2, Chapter 1 General Provisions
- BPC, Division 2, Chapter 9 Pharmacy
- California Code of Regulation (CCR), Title 16, Division 17 California State Board of Pharmacy
- CCR, Title 22, Division 5, Chapter 1 General Acute Care Hospitals
- Civil Code, Division 1, Part 2.6, Chapter 2 Disclosure of Medical Information by Providers
- Code of Federal Regulations (CFR), Title 16, Chapter II, Subchapter E, Part 1700 Poison Prevention Packaging
- CFR, Title 21, Chapter I, Subchapter C, Part 201, Subpart B Labeling Requirements for Prescription Drugs and/or Insulin
- CFR, Title 21, Chapter I, Subchapter C, Part 208 Medication Guides for Prescription Drug Products
- CFR, Title 21, Chapter I, Subchapter C, Part 290 Controlled Drugs
- CFR, Title 21, Chapter I, Subchapter D, Part 310, Subpart E Requirements for Specific New Drugs and Devices
- CFR, Title 21, Chapter II Drug Enforcement Administration, Department of Justice
- Health and Safety Code (HSC), Division 10 Uniform Controlled Substances Act
- HSC, Division 104, Part 5 (Sherman Food, Drug, and Cosmetics Law), Chapter 2 Administration
- HSC, Division 116 Surplus Medication Collection and Distribution
- United States Code (USC), Title 15, Chapter 39A Special Packaging of Household Substances for Protection of Children
- USC, Title 21, Chapter 9, Subchapter V, Part H Pharmaceutical Distribution Supply Chain (Drug Supply Chain Security Act)