

# PHARMACY TECH PTO



(4) Emperating you need to know about passing the 1710.

#### CERTIFICATION



#### The PTCE

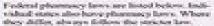
The PTCE is computer based and lasts two-bours, it has 90multiple-choice exections that must be asserted in the first. hour and fifty minutes. The PTCB offers a "blueprint," with an approximate percentage of questions related to specific content areas. However, there are several versions of the FFCE exam, and students taking the same test in the same location will receive different versions of the exam. Some may contain more or low of a porticular content area than another. Once you have period the PTCE, you are estitled to use the CPhT designation. You must recertify with the PTCB every two years and complete 20 hours of continuing

education (CE) per two-year recontilication cycle.

#### Liconoure/Registration by State

In addition to the national certification, you may be expained to become certified or resistered with year state characters. bound. Each state has different requirements for pharmacy. technicisms, and some do not require either national certifigurious or state registration.

#### LAWS



EX: Some medications are considered a controlled substance under state laws, but, more Mandowell.

## Federal Food, Drug & Cosmotic Act of 1938 (FOCA)

- Effectively enested the Food and Drug Administration (FDA), giving them the propossibility to appears all new drags and assume equity standards by endocrine the rules against adultoration and misbranding. Adultoration includes contamination or failure to seect postly and appeliets standards, Masheunding implodes false, insecurity, improper, or confusing labeling or if the product is humaful when used according to the label discritions.
- . The FDCA pives the FDA the power to enforce revalls if any product is in violation of these rules. These recells may opens at the wholevaler, retail, or comconnect level. There are three levels of BIDAN population
  - Class B: For risk of Surrious subverse health extraoperaces or death."
  - Class III: For risk of "temperaturily or modically reversible adverse health consequences" or how-probability of serious adverse besitis consequences. Chass III: For products unlikely to have
  - adverse bealth consequences.

#### Health insurance Portability & Accountability Act of 1996 (HIPAA)

HIPAA is for reaching and regulates the way. health care personnel handle Protected Health Information (PRR), HIPSAA defines PRR as any individually identifiable books information. Under HIPSAA, phomoscies must:

- Ensure the confidentiality of PBR. This includes all manner of transmission, whether written, electronic, or oral (potient counseling, phone calls with providers, etc.). Under HIPAA, phoronacy stuff. may not disclose any PHI to anyone
  - other than the patient, including apoinen and relatives unless the nations has principal extensions commercial.
    - Train pharmacy staff on HIPAA. perculations and periodically retrainor update training.
- · Protect against possible branches with MANAGERY STATESTAND Dispose of PHII in accordance with:
- HIRAA (through measures such as incinexplained or otherodiffence's

- Appoint a designated privacy officer.
- Inform patients of their privacy rights and how the pharmacy handles their in-Commence of the last
- Provisée autients with their PHII records. id appropriately.

Most importantly, any breach must be reported in a timely manner to the Department. of Health and Human Services Office for Civil Rights. If the breach affects fewer than 500 individuals, it must be reported within 60 days of the end of the calendar year. If it affects more than 500 individuals, it must be reported within 60 days of the Service St.

#### Occupational Safety & Health Act of 1970

The Occupational Safety and Health Act of 1970 counted the Occupational Sofety and Health Administration (OSHA) to ensure the soliety of workers nationwisk. As it nelistes to plantaccy, OSHA requires workers to be trained in safety measures, safety observe to be available for all barrandons. substances, and personal protective compmost (PPE) to be more/unificed when exprovince to hazandous substances in presubtic.

#### Controlled Substance Act of 1970

This not erested formal scheduling of drags with the potential for abuse and dependency as controlled substances. It led to the exection of the Drug Enforcement Adestimistration (DEA) in 1973 to help with regulatory policeposent.

#### DEA Number Calculations

A DEA number is required to prescribe controlled substances except for he military practitioners and employees in public-hooth services, prisons, and cortain long-term care facilities. Each DEA number has its own. validation within its digits. Each DEA purpber consists of two fetters and serion digits, enablism you to manually validate a DEA. manuface. Higgs 's horse:

- . The first letter corresponds to the presombor 's registration type.
- This presented better in the first better of the. proposatelloger's light married.
- The lat had and his monthers are added. together, and the 2nd, 4th, and 6th letters. are added to exilter and multiplied by 2. Add the two resulting figures together, and the final digit should match the 7th digit, also known as the check digit.

EX: Dr. James Beyont words in a proscription with a DEA number of

#### AUDIO STREET,

6 - 9 - 4 - 19

医中枢中毒一种化类一种

total or that or W.S.

Since the II in the DEA number marches the "15" in Deviant, and the 7 in 37 metches the check digit, this is a potentially washed EDE A moundoor.

#### Schodules (I-V)

Controlled substances are enterorized into five different schodules, according to their potential for always and decondency:

Netherhole II: No accepted medical use high alreasy potential.

EX: Ellegal design such as LSD and benoise

Note: Cocaine is part schedule Libert. schedule II, although it is rarely used mendicative.

Schodule Hit High above percential. According to the DEA, me of schedule II drugs potentially leads to "severe psychologicall or physical depondence.

EX1 ADHID design such as Ritalia, Concerts, and Addorall; opioid analyseses: such as exceptione and oxygodone; and synthetic opioids soull pe feetare li

Schedule III: Moderate abuse and dependency percential.

EX: Codeine-containing products (less. these 90 may's and testoodyroog-

Schardule IV: Low above and dependency protection.

EX: Benrodiscopines and sleep aids with

Schedule Vt Lowest abuse and dependisplays pertopolish.

EX: Codeine-containing cough syrups. Lordon, Vinnesott

#### Polson Prevention Packaging Act 94 1970

This not requires child safety caps on all disposand prescriptions. Easy open caps more by used of requested, and a record must he kept of the request.

#### Drug Litting Act of 1972

This act established rules for all drug munadactioners to supply a current drug flot to the FDA: Each drug is assigned its own NDC quational drug code). The NDC is

unique to the manufacturer and puckage size; thus, the same medication from difforest manufacturers (e.g., generics) will each have their own NDC. An NDC is seeally booken down into three sections.

- · The first section refers to the manufactumor out reproclamator.
- The second section relies to the specific. drug itself (including strength and form).
- The third section refers to the puckaging.

## Prescription Drug Marketing Act of 1987

- Receives who knowled distribution is be lineweed.
- Sumpley of prescription medication man only be given to liquinoid presonings.
- Expected prescription drugs may not be reimpostod.

## Omnibus Budget Reconciliation Act of 1990 (OBRA-90)

OBSKA was a widespread federal deficit redoction budget bill, the phormacy implications of which initially only pertained to Medicaid recipients: Hirwever, as it was uphe individual states to determine how to enforce OBIRA, most expanded the not to inclosic all potients. Under OBRA, three major noquirements are pertinent for phormacy:

- Phorosociete must office connecting to all. potients, documenting patient refusals. for counsel. (Codor no circumstances should a techniques commed patients.)
- Phoenacists most parties a "prospective" drug utilization periors" (DCR) prior to filling each prescription, which evaluates thorney for potential problems (e.g., interactions, affergies, therapeutic duplication, and other adverse effects).
- Record-beoping mandates: Phormacies. must keep patient profiles of medications. dispensed for two years encode.

#### Combat Mathamphatamina Epidemic Act of 2005

- · Requires pharmacies to limit sales of server-they-counter medications contain prevalorphedrine or ephodrine to 3.6 grams per purchaser per day and no more than 7.5 grams within a 30-day period. Products containing these ingredients must be kept behind the pharmacy counter or in a locked collinet and can only he purchased by contemper over the age of 18 with a valid photo ID.
- Phormacies must maintain a legbook of each repulated psychase, including the customer's signature, address, product purchased quantity purchased, and date and time of the sale. This implock may be physical or electronic and must be maintained for a period of two years.

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