

PUBLIC HEALTH CODE (EXCERPT)
Act 368 of 1978

333.17762 Misbranded prescription.

Sec. 17762.

(1) A prescription drug is considered misbranded unless the manufacturer's label states the name and place of business of the manufacturer of the finished dosage form of a drug and, if different, the name and place of business of the packer or distributor.

(2) As used in this section, "finished dosage form of a drug" means that form of the drug which is or is intended to be dispensed or administered to the patient and does not require further manufacturing or processing other than packaging or labeling, or both.

History: 1978, Act 368, Eff. Sept. 30, 1978

Popular Name: Act 368