ARTICLE 2. CONTROLLED SUBSTANCES

Rule 1. Definitions

856 IAC 2-1-1 Definitions

Authority: IC 35-48-3-1 Affected: IC 4-21.5; IC 35-48-2-1

Sec. 1. Definitions. As used herein, the following terms shall have the meanings specified:

(a) The term "Act" means the Indiana Uniform Controlled Substances Act of 1973. IC 1971, 35-24.1 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended.

(b) The term "basic class" means, as to controlled substances listed in Schedules I and II [856 IAC 2-2-2 and 856 IAC 2-2-3]: (1) Each of the opiates, including its isomers, esters, ethers, salts, and salts of isomers, esters, and ethers whenever the existence of such isomers, esters, ethers, and salts is possible within the specific chemical designation, listed in Section 2.11(b) [856 IAC 2-2-2(b)] of this chapter;

(2) Each of the opium derivatives, including its salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of isomers is possible within the specific chemical designation, listed in Section 2.11(c) [856 IAC 2-2-2(c)] of this part;

(3) Each of the hallucinogenic substances, including its salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of isomers is possible within the specific chemical designation, listed in Section 2.11(d) [856 IAC 2-2-2(d)] of this part;

(4) Each of the following substances whether produced directly or indirectly by extraction from substances of vegetable origin, or independently by means of chemical synthesis, or by a combination of extraction and chemical synthesis:

(i) Opium, including raw opium, opium extracts, opium fluid extracts, powdered opium, granulated opium, deodorized opium and tincture of opium;

- (ii) Apomorphine;
- (iii) Codeine;
- (iv) Ethylmorphine;
- (v) Hydrocodone;
- (vi) Hydromorphone;
- (vii) Metopon;
- (viii) Morphine;
- (ix) Oxycodone;
- (x) Oxymorphone;
- (xi) Thebaine;
- (xii) Mixed alkaloids of opium listed in Section 2.12(b)(2) [856 IAC 2-2-3(b)(2)] of this part;
- (xiii) Cocaine; and
- (xiv) Ecgonine;

(5) Each of the opiates, including its isomers, esters, ethers, salts, and salts of isomers, esters, and ethers whenever the existence of such isomers, esters, ethers and salts is possible within the specific chemical designation, listed in Section 2.12(c) [856 IAC 2-2-3(c)] as amended, of this part;

(6) Methamphetamine, including salts, isomers, and salts of isomers.

(7) Amphetamine, its salts, optical isomers and salts of its optical isomers;

- (8) Phenmetrazine and its salts; and
- (9) Methylphenidate.

(c) The term "Administration" means the Drug Enforcement Administration, formerly the Bureau of Narcotics and Dangerous Drugs.

(d) The term "agent" means an authorized person who acts on behalf of or at the direction of a manufacturer, distributor, or dispenser. It does not include a common or contract carrier, public warehouseman, or employee of the carrier or warehouseman.

(e) The term "controlled premises" means-

(1) Places where original or other records or documents required under the Act [IC 35-48] are kept or required to be kept, and

(2) Places including factories, warehouses, or other establishments, conveyances, where persons registered under the Act *[IC 35-48]* or exempted from registration under the Act *[IC 35-48]* may lawfully hold, manufacture, or distribute, dispense, administer, or otherwise dispose of controlled substances.

(f) The term "Administrator" means the Director of the Federal Drug Enforcement Administration who has been delegated authority under the Controlled Substances Act of 1970 (84 Stat. 1242; 21 U.S.C. 801) by the Attorney General of the United States (28 C.F.R. 0.100), as amended.

(g) The term "hearing" means any hearing held pursuant to the provisions of IC 1971, 4-22-1 through 4-22-1-30 [*Repealed by P.L.18-1986, SECTION 2. See IC 4-21.5.*] as amended and 4-22-2, for the purpose of granting, denying, or revoking, or suspending a registrant or application for registrant or a hearing amending these rules pursuant to IC 1971, 35-24.1 [*Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.*] as amended.

(h) The term "individual practitioner" means a physician, dentist, veterinarian, or other individual licensed, registered or otherwise permitted by the State of Indiana or the United States, to dispense a controlled substance in the course of professional practice, but does not include a pharmacist, a pharmacy, or an institutional practitioner.

(i) The term "institutional practitioner" means a hospital or other person (other than an individual) licensed, registered, or otherwise permitted by the state of Indiana or the United States, to dispense a controlled substance in the course of practice, but does not include a pharmacy.

(j) The term "person" includes any individual, corporation, government or governmental subdivision or agency, business, trust partnership, association or other legal entity.

(k) The term "pharmacist" means any practitioner licensed as a pharmacist by the State of Indiana to dispense controlled substances and shall include pharmacist interns licensed by the State of Indiana, to dispense controlled substances under the supervision of a pharmacist licensed by the State of Indiana.

(1) The term "prescription" means an order for medication which is dispensed to or for an ultimate user but does not include an order for medication which is dispensed for immediate administration to the ultimate user. (e.g., an order to dispense a drug to a bed patient for immediate administration in a hospital is not a prescription).

(m) The terms "register" and "registration" refers only to registration required and permitted by IC 1971, 35-24.1-3-2 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended.

(n) The term "registrant" means any person who is registered or exempted from registration pursuant to IC 1971, 35-24.1-3-2 *[Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.]* as amended.

(o) Any term not defined in this section shall have the definition set forth in IC 1971, 35-24.1-1-1 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended.

(Indiana Board of Pharmacy; Reg 28, Ch I, Sec 1.01; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-1-2 Controlled substances advisory committee; function; hearings

Authority: IC 35-48-3-1

Affected: IC 4-21.5; IC 35-48-2-1

Sec. 2. Function. The Controlled Substances Advisory Committee shall serve as a consultative and advisory body to the Board in all matters relative to additions, deletions and transfers of substances to or among schedules of control established by IC 1971, 35-24.1 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended.

In addition, the advisory committee may, as representatives of the Board, conduct hearings regarding control of substances, and it shall, as representatives of the Board, conduct hearings and make recommended findings in matters affecting the denial, suspension, or revocation of registrations. All adjudicatory hearings shall be conducted in a manner consistent with the provision of IC 1971, 35-24.1-3-4 through IC 1971, 35-24.1-3-5 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48], and IC 1971, 4-22-1 [Repealed by P.L.18-1986, SECTION 2. See IC 4-21.5.] as amended. (Indiana Board of Pharmacy; Reg 28, Ch I, Sec 1.11; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-1-3 Meetings; organization

Authority: IC 35-48-3-1 Affected: IC 35-48-2-1 Sec. 3. Meetings and Organization. The controlled substances advisory committee shall meet not later than sixty (60) days after the appointment of their entire membership and thereafter shall meet upon the request of the Board. The committees shall select, from among their members, a chairman, vice-chairman, and secretary who shall serve terms of one year from the date of selection. In any case in which a committee officer shall be unable to serve a full term, the committee shall select another to serve in his own right a full term. (Indiana Board of Pharmacy; Reg 28, Ch I, Sec 1.12; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-1-4	Duties of officers	
Authority:	IC 35-48-3-1	
Affected:	IC 35-48-2-1	

Sec. 4. Duties of Officers. The chairman of the committee, or the vice-chairman in the absence of the chairman, shall preside at all meetings of the committee. In addition, the chairman or his designee shall preside over all hearings conducted by the committee on behalf of the Board.

The secretary of the committee shall be responsible for keeping the minutes of all meetings and he shall further be charged with the responsibility of assuring that a complete and accurate record is made of all hearings conducted before the committee. To this end, he may, with the consent of the Board, arrange for the attendance of such stenographers or court reporters as are necessary for the recording of such hearings. (Indiana Board of Pharmacy; Reg 28, Ch I, Sec 1.13; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-1-5 Rulemaking powers

Authority: IC 35-48-3-1 Affected: IC 4-22-2; IC 35-48-2-1

Sec. 5. Rules of Conduct. The advisory committee may, with the approval of the Board, make such other rules regulating its conduct and procedure as are necessary and proper for the orderly conduct of its business.

All such rules, when they may affect procedure or substance of matters which may come before the Board for adjudication, after promulgation in accordance with IC 1971, 4-22-2 as amended, shall be in writing and shall be made available upon request to parties appearing before the committee. (Indiana Board of Pharmacy; Reg 28, Ch I, Sec 1.14; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-1-6 Recommendations and findings

Authority: IC 35-48-3-1 Affected: IC 35-48-2-1

Sec. 6. Recommendations and findings. Recommendations and findings to be in writing. Whenever, in the discharge of its duties, the advisory committee shall be required to make recommendations or findings upon matters heard before the committee, such recommendations to the Board shall be in writing and shall include a summary of relevant evidence, opinions, and laws upon which such recommendations or findings are based. (Indiana Board of Pharmacy; Reg 28, Ch I, Sec 1.15; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

Rule 2. Controlled Substances Code Number–Schedules I through IV

856 IAC 2-2-1 Controlled substances code numbers

Authority: IC 35-48-3-1 Affected: IC 35-48-3-1

Sec. 1. Controlled Substances Code Number. (a) Each controlled substance, or basic class thereof, listed in Schedules I through IV [856 IAC 2-2-2 – 856 IAC 2-2-5] has been assigned a "Controlled Substances Code Number" for purposes of identification of the substances or class on certain Certificates of Registration issued by the Indiana State Board of Pharmacy pursuant to Section 3.42

[856 IAC 2-3-19] of the Chapter. Certain applicants for registration must include the appropriate numbers on the application as required in Section 3.32(d) [856 IAC 2-3-13(d)] of this Chapter.

(b) Except as stated in paragraph (a) of this section, no applicant or registrant is required to use the Controlled Substances Code Number for any purpose. (Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.01; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-2 Schedule I

Authority: IC 35-48-2-14; IC 35-48-3-1 Affected: IC 35-48-2-4

Sec. 2. (a) Schedule I shall consist of the drugs and other substances, by whatever official name, common or usual name, chemical name, or brand name designated, listed in this section. Each drug or substance has been assigned the controlled substances code number set forth opposite it.

(b) Opiates. Unless specifically excepted or unless listed in another schedule, any of the following opiates, including its isomers, esters, ethers, salts, and salts of isomers, esters, and ethers whenever the existence of such isomers, esters, ethers, and salts is possible within the specific chemical designation:

(1) Acetylmethadol	9601
(2) Allylprodine	
(3) Alphacetylmethadol (except levo-alphalmethadol also known as levo-alpha-acetylmethadol, levomethadyl acetate	
LÁAM)	
(4) Alphameprodine	9604
(5) Alphamethadol	
(6) Benzethidine	
(7) Betacetylmethadol	9607
(8) Betameprodine	9608
(9) Betamethadol	9609
(10) Betaprodine	9611
(11) Clonitazene	9612
(12) Dextromoramide	
(13) Dextrorphan	9614
(14) Diampromide	9615
(15) Diethylthiambutene) 616
(16) Difenoxin) 168
(17) Dimenoxadol) 617
(18) Dimepheptanol) 618
(19) Dimethylthiambutene) 619
(20) Dioxaphetyl butyrate) 621
(21) Dipipanone	
(22) Ethylmethylthiambutene	9623
(23) Etonitazene	
(24) Etoxeridine	
(25) Furethidine	
(26) Hydroxypethidine	
(27) Ketobemidone	
(28) Levomoramide	
(29) Levophenacylmorphan	
(30) Morpheridine	
(31) Noracymethadol	
(32) Norlevorphanol	
(33) Normethadone	
(34) Norpipanone) 636

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(35) Phenadoxone	
(36) Phenampromide	
(37) Phenomorphan	
(38) Phenoperidine	
(39) Piritramide	
(40) Proheptazine	
(41) Properidine	
(42) Propiram	
(43) Racemoramide	
(44) Trimeperidine	
(c) Opium derivatives. Unless specifically excepted or unless listed in another schedule, any of the following c	
derivatives, its salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of isomers is post	ssible
within the specific chemical designation:	
(1) Acetrophine	9319
(2) Acetyldihydrocodeine	
(3) Benzylmorphine	9052
(4) Codeine methylbromide	9070
(5) Codeine-N-Oxide	
(6) Cyprenorphine	
(7) Desomorphine	
(8) Dihydromorphine	
(9) Drotebanol	
(10) Etorphine (Except Hydrochloride Salt)	
(11) Heroin	
(12) Hydromorphinol	
(13) Methyldesorphine	
(14) Methylidihydromorphine	
(15) Morphine methylbromide	
(16) Morphine methylsufonate	
(17) Morphine-N-Oxide	
(17) Worphine (18) Myrophine	
(19) Nicocodeine	
(1) Nicocodenie (20) Nicomorphine	
(20) Notonioiphine	
(22) Pholcodine	
(22) Thoreon	
(d) Hallucinogenic substances. Unless specifically excepted or unless listed in another schedule, any material, comp	
mixture, or preparation, which contains any quantity of the following hallucinogenic substances, or which contains any of its	
isomers, and salts of isomers, whenever the existence of such salts, isomers, and salts of isomers is possible within the sp	
chemical designation (for purposes of this subsection only, "isomer" includes the optical, position, and geometric isomers):	ecific
(1) 4-Bromo-2, 5-Dimethoxyamphetamine	7201
Some trade or other names:	/ 391
4-Bromo-2, 5-Dimethoxy-a-methylphenethylamine; 4-Bromo-2, 5-DMA	7206
(2) 2, 5-Dimethoxyamphetamine	/390
Some trade or other names:	
2, 5-Dimethoxy-a-methylphenethylamine; 2,5-DMA	7 4 1 1
(3) 4-Methoxyamphetamine	/411
Some trade or other names:	
4-Methoxy-a-methylphenethylamine: Paramethoxyamphetamine: PMA	7401
(4) 5-methoxy-3, 4-methylenedioxy amphetamine	
(5) 4-methyl-2, 5-dimethoxyamphetamine	/395

Some trade and other names:
4-methyl-2,5-dimethoxy-a-methylphenethylamine: "DOM"; and "STP".
(6) 3, 4-methylenedioxy amphetamine
(7) 3, 4, 5-trimethoxy amphetamine
(8) Bufotenine
Some trade and other names:
3-(B-Dimethylaminoethyl)-5-hydroxyindol; 3-(2-Dimethylaminoethyl)-5-indolo; N, N-dimenthylserotonin; 5-
hydroxy-N, N-dimethyltryptamine; mappine.
(9) Diethyltryptamine
Some trade and other names: N, N-Diethyltryptamine, DET.
(10) Dimethyltryptamine
Some trade or other names: DMT
(11) Ibogaine
Some trade and other names:
7-Ethyl-6, 6a, 7, 8, 9, 10, 12, 13-octahydro-2-methoxy-6, 9-methano-5H-pyrido (1', 2': 1, 2) azepino 4, 5-b)
indole; tabernanthe iboga.
(12) Lysergic acid diethylamide
(13) Marihuana
(14) Mescaline
(15) Peyote
Meaning all parts of the plant presently classified botanically as Lophophora Williamsii Lemaire, whether growing or
not; the seeds thereof; any extract from any part of such plant; and every compound, manufacture, salt, derivative,
mixture, or preparation of such plant, its seeds, or extracts.
(Interprets 21 U.S.C. 812(c), Schedule I(c) (12))
(16) N-ethyl-3-piperidyl benzilate
(17) N-methyl-3-piperidyl benzilate
(18) Psilocybin
(19) Psilocyn
(20) Tetrahydrocannabinols
Synthetic equivalents of the substances contained in plant, or in the resinous extractives of Cannabis, sp. and/or synthetic
substances, derivatives, and their isomers with similar chemical structure and pharmacological activity such as the
following:
\triangle^1 cis or trans tetrahydrocannabinol, and their optical isomers.
\triangle^6 cis or trans tetrahydrocannabinol and their optical isomers.
$\triangle^{3,4}$ cis or trans tetrahydrocannabinol, and its optical isomers.
(Since nomenclature of these substances is not internationally standardized, compounds of these structures, regardless
of numerical designation of atomic positions are covered.)
(21) Thiophene Analog of Phencyclidine
Some trade or other names:
1-(1-(2-thienyl) cyclohexyl) piperidine); 2-Thienyl Analog of Phencyclidine, TPCP.
(e) Depressants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or
preparation which contains any quantity of the following substances having a depressant effect on the central nervous system,
including its salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of isomers is possible within
the specific chemical designation:
(1) Mecloqualone
(Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.11; filed Jul 9, 1974, 9:29 a.m.: Unpublished; filed Jun 9, 1977, 8:55 a.m.:
Unpublished; filed May 31, 1994, 5:00 p.m.: 17 IR 2335; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct

4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-3 Schedule II

Authority: IC 35-48-2-14; IC 35-48-3-1 Affected: IC 35-48-2-6

Sec. 3. (a) Schedule II shall consist of the drugs and other substances, by whatever official name, common or usual name, chemical name, or brand name designated, listed in this section. Each drug or substance has been assigned the controlled substances code number set forth opposite it.

(b) Substances, vegetable origin, or chemical synthesis. Unless specifically excepted or unless listed in another schedule, any of the following substances whether produced directly or indirectly by extraction from substances of vegetable origin, or independently by means of chemical synthesis, or by a combination of extraction and chemical synthesis:

(1) Opium and opiate, and any salt, compound, derivative, or preparation of opium or opiate, excluding naloxone and its salts and naltrexone and its salts but including the following:

(A) Raw opium	9600
(B) Opium extracts	9610
(C) Opium fluid extracts	9620
(D) Powdered opium	9639
(E) Granulated opium	9640
(F) Tincture of opium	9630
(F) Tincture of opium	9030
(H) Codeine	9050
(I) Ethylmorphine	
(J) Etorphine hydrochloride	9059
(K) Hydrocodone	9193
(L) Hydromorphone	9194
(M) Metopon	9260
(N) Morphine	9300
(O) Oxycodone	9143
	9652
(O) Thebaine	9333

(2) Any salt, compound, derivative, or preparation thereof which is chemically equivalent or identical with any of the substances referred to in subdivision (1), except that these substances shall not include the isoquinoline alkaloids of opium.(3) Opium poppy and poppy straw (9650).

(4) Coca Leaves (9040) and salt, compound, derivative, or preparation of coca leaves, and any salt, compound, derivative, or preparation thereof which is chemically equivalent or identical with any of these substances, except that the substances shall not include decocainized coca leaves or extraction of coca leaves, which extractions do not contain cocaine (9041) or ecgonine (9180).

(5) Concentrate of poppy straw (the crude extract of poppy straw in either liquid, solid, or powder form which contains the phenanthrine alkaloids of the opium poppy) 9670.

(c) Opiates. Unless specifically excepted or unless listed in another schedule, any of the following opiates, including its isomers, esters, ethers, salts, and salts of isomers, esters, and ethers whenever the existence of such isomers, esters, ethers, and salts is possible within the specific chemical designation:

(1) Alphaprodine	9010
(2) Anileridine	9020
(3) Benzitramide	
(4) Dihydrocodeine	9120
(5) Diphenoxylate	9170
(6) Fentanyl	
(7) Isomethadone	9226
(8) Levo-alphacetylmethadol	9648
Some trade and other names:	

levo-alpha-acetylmethadol, levomethadyl acetate, or LAAM.

(9) Levomethorphan	9210
(10) Levorphanol	9220
(11) Metazocine	9240
(12) Methadone	9250
(13) Methadone-Intermediate, 4-cyano-2-dimethylamino-4, 4-dyphenyl butane	9254
(14) Moramide-Intermediate, 2-methyl-3-morpholino-1, 1-diphenyl-propane-carboxylic acid	
(15) Pethidine	9230
(16) Pethidine-Intermediate-A,4-cyano-1- methyl-4-phenylpiperidine	9232
(17) Pethidine-Intermediate-B,ethyl-4-phenylpiperidine-4-carboxylate	9233
(18) Pethidine-Intermediate-C,1-methyl-4- phenylpiperidine-4-carboxylic acid	9234
(19) Phenazocine	9715
(20) Piminodine	9730
(21) Racemethorphan	9732
(22) Racemorphan	
(d) Stimulants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixtu	
preparation which contains any quantity of the following substances having a stimulant effect on the central nervous system	
(1) Amphetamine, its salts, optical isomers, and salts of its optical isomers	
(2) Methamphetamine, including its salts, isomers, and salts of isomers	
(3) Phenmetrazine and its salts	
(4) Methylphenidate	
(e) Depressants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixtu	
preparation which contains any quantity of the following substances having a depressant effect on the central nervous sy	
including its salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of isomers is possible	within
the specific chemical designation:	
(1) Methaqualone	
(2) Amobarbital	
(3) Secobarbital	
(4) Pentobarbital	
(Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.12; filed Jul 9, 1974, 9:29 a.m.: Unpublished; filed Jun 9, 1977, 8:55	a.m.:

(Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.12; filed Jul 9, 1974, 9:29 a.m.: Unpublished; filed Jun 9, 1977, 8:55 a.m.: Unpublished; filed May 31, 1994, 5:00 p.m.: 17 IR 2336; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-4 Schedule III

Authority:	IC 35-48-3-1
Affected:	IC 35-48-2-8

Sec. 4. (a) Schedule III shall consist of the drugs and other substances, by whatever official name, common or usual name, chemical name, or brand name designated listed in this section. Each drug or substance has been assigned the Controlled Substances Code Number set forth opposite it.

(b) Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation that contains any quantity of the following substances having a stimulant effect on the central nervous system, including its salts, isomers (whether optical, position, or geometric), and salts of such isomers whenever the existence of such salts, isomers, and salts of isomers is possible within the specific chemical designation:

(1) Those compounds, mixtures, or preparations in dosage unit form containing any stimulant substances listed in Sche	edule
II, which compounds, mixtures, or preparations were listed on August 25, 1971, as excepted compounds under 1308.32	2, and
any other drug of the quantitative composition shown in that list for those drugs or that is the same, except that it conta	ains a
lesser quantity of controlled substances	1405
(2) Benzphetamine	1228
(3) Chlorphentermine	1645
(4) Clortermine	1647
(5) Mazindol	1605

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(6) Phendimetrazine
(c) Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation that
contains any quantity of the following substances having a depressant effect on the central nervous system:
(1) Any compound, mixture, or preparation containing:
(A) Amobarbital
(B) Secobarbital
(C) Pentobarbital
or any salt thereof and one (1) or more other active medicinal ingredient that are not listed in any schedule.
(2) Any suppository dosage form containing:
(A) Amobarbital
(B) Secobarbital
(C) Pentobarbital
or any salt of any of these drugs and approved by the Food and Drug Administration for marketing only as a suppository.
(3) Any substance that contains any quantity of a derivative of barbituric acid or any salt thereof
(4) Chlorhexadol
(5) Ketamine, its salts, isomers, and salts of isomers
Some other names for ketamine: (-2(2-chlorophenyl) -2- (methylamino) - cyclohexanone
(6) Lysergic acid
(7) Lysergic acid amide
(8) Methyprylon
(9) Sulfondiethylmethane
(10) Sulfonethylmethane
(11) Sulfonmethane 2610
(d) Nalorphine (a narcotic drug)
(e) Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation
containing limited quantities of any of the following narcotic drugs, or any salts thereof:
(1) Not more than one and eight-tenths (1.8) grams of codeine, per one hundred (100) milliliters or not more than ninety (90) milligrams per dosage unit, with an equal or greater quantity of an isoquinoline alkaloid of opium
(2) Not more than one and eight-tenths (1.8) grams of codeine, per one hundred (100) milliliters or not more than ninety (90)
milligrams per dosage unit, with one (1) or more active, nonnarcotic ingredients in recognized therapeutic amounts . 9804
(3) Not more than three hundred (300) milligrams of dihydrocodeinone, per one hundred (100) milliliters or not more than
fifteen (15) milligrams per dosage unit, with a fourfold or greater quantity of an isoquinoline alkaloid of opium 9805
(4) Not more than three hundred (300) milligrams of dihydrocodeinone, per one hundred (100) milliliters or not more than
fifteen (15) milligrams per dosage unit, with one (1) or more active, nonnarcotic ingredients, in recognized therapeutic amounts
9806
(5) Not more than one and eight-tenths (1.8) grams of dihydrocodeine, per one hundred (100) milliliters or not more than ninety
(90) milligrams per dosage unit, with one (1) or more active, nonnarcotic ingredients in recognized therapeutic amoun 9807
(6) Not more than three hundred (300) milligrams of ethylmorphine, per one hundred (100) milliliters or not more than fifteen
(15) milligrams per dosage unit, with one (1) or more active, nonnarcotic ingredients in recognized therapeutic amoun@808
(7) Not more than five hundred (500) milligrams of opium per one hundred (100) milliliters or per one hundred (100) grams
or not more than twenty-five (25) milligrams per dosage unit, with one (1) or more active, nonnarcotic ingredients in
recognized therapeutic amounts
(8) Not more than fifty (50) milligrams of morphine, per one hundred (100) milliliters or per one hundred (100) grams with
one (1) or more active, nonnarcotic ingredients in recognized therapeutic amounts
(f) Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation
containing any quantity of anabolic steroids, including its salts, isomers, and salts of isomers whenever the existence of such salts
of isomers is possible within the specific chemical designation
(g) For hallucinogenic substances, dronabinol (synthetic) in sesame oil and encapsulated in a soft gelatin capsule in a United
States Food and Drug Administration-approved drug product
(Some other names for dronabinol: (6aR-trans) - 6a, 7, 8, 10a - tetrahydro-6,6,9 - trimethyl - 3-pentyl- 6H - dibenzo[b,d]pyrano-1-ol,
or (1) \triangle^9 - (trans) - tetrahydrocan-nabinol.) (Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.13; filed Jul 9, 1974, 9:29 a.m.:

Unpublished; filed Jun 9, 1977, 8:55 a.m.: Unpublished; filed May 26, 2000, 8:52 a.m.: 23 IR 2503; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-	-5 Scł	edule IV
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Authority: IC 35-48-3-1 Affected: IC 35-48-2-10

Sec. 5. (a) Schedule IV shall consist of the drugs and other substances, by whatever official name, common or usual name, chemical name, or brand name designated, listed in this section. Each drug or substance has been assigned the Controlled Substances Code Number set forth opposite it.

(b) Depressants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substances, including its salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of isomers is possible within the specific chemical designation:

(1) Barbital	2145
(2) Chloral betaine	2460
(3) Chloral hydrate	2465
(4) Chlordiazepoxide	2744
(5) Clonazepam	2737
(6) Clorazepate	2768
(7) Diazepam	2765
(8) Ethchlorvynol	2540
(9) Ethinamate	2545
(10) Flurazepam	2767
(11) Mebutamate	2800
(12) Meprobamate	2820
(13) Methohexital	2264
(14) Methylphenobarbital	2250
(15) Oxazepam	2835
(16) Paraldehyde	2585
(17) Petrichloral	2591
(18) Phenobarbital	2285
(a) Earflyraming Any material compound mixture or propagation which contains any quantity of the following such	atomaga

(c) Fenfluramine. Any material, compound, mixture, or preparation which contains any quantity of the following substances, including its salts, isomers (whether optical, position, or geometric), and salts of such isomers, whenever the existence of such salts, isomers, and salts of isomers is possible:

(d) Stimulants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substances having a stimulant effect on the central nervous system, including its salts, isomers (whether optical, position, or geometric), and salts of such isomers whenever the existence of such salts, isomers is possible within the specific chemical designation:

(1) Diethylpropion	1608
(2) Phentermine	1640
(3) Pemoline (including organometallic complexes and chelates thereof)	1530

(Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.14; filed Jul 9, 1974, 9:29 am: Unpublished; filed Jun 9, 1977, 8:55 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-6 Schedule V

Authority: IC 35-48-3-1 Affected: IC 35-48-2-12

Sec. 6. (a) Schedule V shall consist of the drugs and other substances, by whatever official name, common or usual name,

chemical name, or brand name designated, listed in this section.

(b) Narcotic drugs containing nonnarcotic active medicinal ingredients. Any compound, mixture, or preparation containing any of the following limited quantities of narcotic drugs or salts thereof, which shall include one or more nonnarcotic active medicinal ingredients in sufficient proportion to confer upon the compound, mixture, or preparation valuable medicinal qualities other than those possessed by the narcotic drug alone:

(1) Not more than 200 milligrams of codeine, per 100 milliliters or per 100 grams.

(2) Not more than 100 milligrams of dihydrocodeine, per 100 milliliters or per 100 grams.

(3) Not more than 100 milligrams of ethylmorphine, per 100 milliliters or per 100 grams.

(4) Not more than 2.5 milligrams of diphenoxylate and not less than 25 micrograms of atropine sulfate per dosage unit.

(5) Not more than 100 milligrams of opium per 100 milliliters or per 100 grams.

(Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.15; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-7 Application for exception of stimulant or depression compound; revocation

Authority: IC 35-48-3-1 Affected: IC 35-48-2-1

Sec. 7. Application for exception of a stimulant or depressant compound. (a) Any person seeking to have any compound, mixture, or preparation containing any depressant or stimulant substance listed in Chapter 2, Section 2.13(b) or (c) [856 IAC 2-2-4(b) or (c)] or in Section 2.14 [856 IAC 2-2-5] excepted from the application of all or any part of the Act [IC 35-48] pursuant to IC 1971, 35-24.1-2-8(e) or 10(c) [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended, may apply to the Indiana Board of Pharmacy for such exception.

(b) An application for an exception under this section shall be approved by the Board, provided that such application shall contain a copy of the Administration's approval for such request for an exception from the Federal Controlled Substances Act.

(c) Within a reasonable period of time after the receipt of an application for an exception under this section, the Indiana State Board of Pharmacy shall notify the applicant of its acceptance or non-acceptance of the application, and if not accepted, the reason therefor. The Indiana State Board of Pharmacy need not accept an application for filing if any of the requirements prescribed in paragraph (b) of this section is lacking or is not set forth so as to be readily understood. If the applicant desires, he may amend the application to meet the requirements of paragraph (b) of this section.

(d) The Indiana Board of Pharmacy may at any time revoke any exception granted pursuant to IC 1971, 35-24.1-2-8(e) or 10(c) [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.], as amended, upon a finding that such exception from the Federal Controlled Substances Act was terminated, suspended or revoked. The revocation of the exception granted under this Act [IC 35-48] shall become effective upon the Board's notifying the person to whom such exception was granted by certified mail of such revocation. (Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.21; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-8 Excepted stimulant or depressant compounds

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Authority: IC 35-48-3-1
Affected: IC 35-48-2-8; IC 35-48-2-10
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Sec. 8. The Indiana Board of Pharmacy may except any compound, mixture, or preparation containing any depressant or stimulant substance listed in Section 2.13(b) or (c) [856 IAC 2-2-4(b) or (c)], or in Section 2.14 [856 IAC 2-2-5] from the application of all or any part of the Act pursuant to IC 1971, 35-24.1-2-8(e) or 10(c) [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended, the drugs which were excepted by the Bureau or Administration on April 1, 1973 under section 202(d) of the Federal Controlled Substances Act (21 U.S.C. 812(d)) have been excepted by the Indiana State Board of Pharmacy from the application of IC 1971, 25-24.1-3, 6 and 8 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended, and the application of Section 3.74(d) [856 IAC 2-3-33(d)] (rule) for administrative purposes only. The excepting of these drugs by the Indiana State Board of Pharmacy should not be construed as an adoption or rejection of the criteria by which these drugs were originally excepted. Any deviation from the quantitative composition of any of the listed drugs shall require a petition for exception in order for that drug to be excepted.

The following is a list of the excepted stimulant or depressant compounds under these regulations [856 IAC 2-2].

EXCEPTED PRESCRIPTION DRUGS

Trade name or other designation	Composition	Manufacturer or supplier
A.E.A	Tablet : Amobarbital, 25 mg. ; aminophylline,	Haack Laboratories, Inc.
Alased	120 mg.; ephedrine hydrochloride, 25 mg. Tablet: Phenobarbital, 16.2 mg.; homatropine methylbronide, 3.6 mg.; aluminum hydroxide gel, dried, 7½ gr.; magnesium trisilicate,	Norgine Laboratories, Inc.
Alcitex	21/2 gr. Tablet: Phenobarbital, 1/6 gr.; atropine sulfate, 1/5000 gr.; calcium carbonate, 31/2 gr.; mag- nesium carbonate, 21/2 gr.; cerium oxalate,	Paul B. Elder Co., Inc.
Algoson	1/2 gr. Tablet: Butabarbital sodium, 7.5 mg.; aceta- minophen, 800 mg.	McNeil Laboratories, Inc.
Albydrox	Tablet: Phenobsrbital, 1/2 gr.; aluminum hy- droxide, 5 gr.; atropine sulfate, 1/300 gr.	Physicians Supply.
Alkasans	Tablet Phenoberbital 2.0 mg · stroning sul-	P. J. Noyes Co.
Alsical	 fate, 0.06 mg.; kaolin-alumina gel, 500 mg. Powder (60 gr.): Phenobarbital, ½ gr.; bella- donna extract, ¼ gr.; calcium carbonate, 24 gr.; magnesium trialicate, 15 gr.; mag- nesium oxide, 10 gr.; aluminum hyroxide 	Dorsey Laboratories.
Alubelap	gel, dried. 10 gr. Tablet: Phenobarbital, 8 mg.; aluminum hy- droxide gel, dried, 2300 mg.; belladonna ex- tract, 4 mg.	Haack Laboratories, Inc.
Aludrox SA suspension	Suspension (5 cc.): Butabarbital, 3 mg.; am- butonium bromide, 2.5 mg.	Wyeth Laboratories.
Aludrox SA tablets	Tablet: Butabarbital, 8 mg.; ambutonium bromide, 2.5 mg.	Wyeth Laboratories.
Alu-Mag	Tablet: Phenobarbital, ¼ gr.; aluminum hy- droxide gel, dried, 2¼ gr.; magnesium tri- silicate, 2¼ gr.; belladonna leaf extract,	Norsal Laboratories, Inc.
Alumazen	% gr.	The Zemmer Co.
Aluminum hydroxide, mag- nesium trisilicate, and kaolin with phenobarbital	Tablet: Phenobarbital, % gr.; aluminum hy- droxide, 2 gr.; magnesium trisilicate, 4 gr.;	Buffalo Pharmaceutical Supply Corp.
and atropine sulfate Aminodrox with phenobar- bital	Tablet: Phenobarbital, 15 mg.; aminophylline, 0.1 gm., aluminum hydroxide gel, dried, 0.12	The S. E. Massengill Co.
Aminodrox-forte with phe- nobarbital	200 mg.; aluminum hydroxide gel, dried,	
Aminophylline and amytal	250 mg. Capsule: Amobarbital, 32 mg.; aminophylline, 0.1 gm.	Eli Lilly Co.
Aminophylline with pento- barbital		G. D. Searle & Co.
Aminophylline and pheno- barbital	100 mg.	
Do	Tablet: Phenobarbital, ¼ gr.; aminophylline, 100 mg.	The Blue Line Chemical Co
Aminophylline with pheno- barbital	Tablet: Phenobarbital, 16 mg.; aminophylline,	I-a
Do	Tablet: Phenobarbital, 15 mg.; aminophylline, 100 mg.	G. D. Searle & Co.
Do	Tablet: Phenobarbital, 15 mg.; aminophylline	Do.
Do	200 mg. Tablet: Phenobarbital, 30 mg.; aminophylline, 200 mg.	Do.
Amobarbital and PETN		Meyer Laboratories, Inc.
Ampyrox with butabarbital sodium (AMPYROX)	Tablet: Butabarbital sodium, 15 mg.; scopola-	Paul B. Elder Co., Inc.
	Elixir (5 cc.): Butabarbital sodium, 10 mg.: scopolamine methylnitrate, 1 mg.	Do.

Trade name or other designation	Composition	Manufacturer or supplier
Lmsed (NAP-37)	Tablet: Phenobarbital, ¼ gr.; hyocine hydro- bromide, 0.0072 mg.; atropine sulfate, 0.024	North American Pharma- cal, Inc.
Amendyne	mg.; hyoscyamine hydrobromide, 0.128 mg. Tablet: Phenobarbital, 1/gr.; extract belladonna logna 1/ gr. astrong 1/ gr. astrong 1/	Paul B. Elder Co., Inc.
Antacia No. 3 with pheno- barbital and stropine	leaves, ¼ gr.; aspirin, 5 gr.; caffeine, ¼ gr. Tablet: Phenobarbital, ¼ gr.; stropine sulfate, 1/800 gr.; calcium carbonate, 5 gr.; mag- nesium hydroxide, 5 gr.	Meyers and Co.
Antispasmodic	Tablet (purple): Phenobarbital, 16.2 mg.; Byo- scyamine sulfate, 0.1037 mg.; homatropine	Hydrex Co., Inc.
Antispasmodic-ensyme	Tablet: Phenobarbital, 8.1 mg.; hyosine hydro- sulfate, 0.0055 mg.; homatropine methylbro- mide, 0.2855 mg.; hyoseine hydrobromide, 0.0033 mg.; pancreatin, 100 mg.; pepsin, 150 mg.	Do.
	Tablet or capsule: Phenobarbital, 16 mg.; stro- pine sulfate, 0.324 mg.; colloidal sulfur, 22 mg.	Wm. P. Poythress & Co., Inc.
Aqualin-plus, children	Supportions : Destabashital radium 1/ er + then-	The Wm. A. Webster Co.
	phylline, 1% gr. Suppository: Pentobarbital sodium, % gr.; theo- phylline, 3% gr.	Do.
	Suppository: Pentobarbital sodium, 1½ gr.; theophylling, 7% gr.	Do.
	Suppository : Pentobarbital sodium, % gr. ; theo- phylline, 7% gr.	Do.
	Tablet : Butabarbital, 20 mg.; ephedrine sulfate, 25 mg. : theophylline hydroxide, 130 mg.	
	Tablet: Butabarbital, 16 mg.; aminophylline, 180 mg.; phenylpropanolamine hydrochloride, 25 mg.; chlorpheniramine maleate, 2 mg.; aluminum hydroxide gel, dried, 60 mg.; mag- nesium tridlicate 60 mg.; mag-	
Asperesse, modified with phenobarbital	Tablet: Phenobarbital, 0.008 gm.; acetylsall-	P. J. Noyes Co.
Atropal	Tablet: Phenobarbital, ¼ gr.; atropine sul- fate, 1/300 gr.; magnesium trisilicate, 2¼ gr.; aluminum hydroxide gel, dried, 2½ gr.	
Atrosilita]	Tablet: Phenobarbital, 15 mg.; atropine sul- fate, 0.12 mg.; magnesium trisilicate, 0.5 gm.; saccharine sodium, 0.12 mg.	The Zemmer Co.
Banthine with pheno-	Tablet: Phenobarbital, 15 mg.; methantheline	
	bromide, 50 mg. Tablet: Phenobarbital, 15 mg.; atropine sul- fate, 0.12 mg.	
	Tablet: Phenobarbital, 15 mg.; atropine sul- fate, 0.25 mg.	
	Tahlet: Amobarbital sodium, 20 mg.; hyoscy- amine sulfate, 0.125 mg.; hyoscine hydrobro- mide, 0.007 mg.; homatropine methylbromide, 0.5 mg.	
Barbidonna elixir	Elixir (5 cc.): Phenobarbital, 16 mg.; hyoscy- amine sulfate, 0.1286 mg.; atropine sulfate, 0.0250 mg.; scopolamine hydrobromide, 0.0074 mg.	WOLKS.
	Tablet: Phenobarbital, 16 mg.; hyoscyamine sulfate, 0.1286 mg.; atropine aulfate, 0.0250 mg.; scopolamine hydrobromide, 0.0074 mg.	
Barboma elixir	Elixir (100 cc.) : Phenobarbital, 0.4 gm. ; homa- tropine methylbromide, 33.8 mg.	·
	Tablet : Phenobarbital, ¼ gr. ; homatropine methylbromide, 123 gr.	
Bardase	Tablet or elixir (4 cc.): Phenobarbital, 16. mg.; hyoscyamine sulfate, 0.1 mg.; hyoscin- hydrobromide, 0.007 mg.; atropine, 0.020 mg. Taka-Diastase, 162.0 mg.	e

Truis name or other designation	Competition	Manufacturer or supplier
Ber-Don elixir	Elixir (30 ec.): Phenobarbital, 100 mg.; hyoscy- amine hydrobromide, 0.60 mg.; hyoscine hy- drobromide, 0.042 mg.; atropine sulfate, 0.12 mg.	Warren-Teed Pharmaceut cals, Inc.
Bar-Don tablets	Tablet : Phenobarbital, 16.670 mg.; hyoscyamine hydrobromide, 0.10 mg.; hyoscine hydrobro- mide, 0.007 mg.; atropine sulfate 0.020 mg.	Do.
Belap No	Tablet: Phenobarbital, 8 mg.; belladonna ex- tract, 8 mg.	Haack Laboratories, Inc.
Belap No. 1	Tablet: Phenobarbital, 15 mg.; belladonna ex- tract. 8 mg.	Do.
Belap Ty-Med	methylbromide, 7.5 mg.	Do.
Belladepal	mg.	Do.
	Elixir (15 cc.): Phenobarbital, 15.6 mg.; bella- foliae, 0.075 mg.	
	Elixir (5 cc.): Butabarbital sodium, 20 mg.; tincture belladonna, 0.83 cc.	
	Tablet : Phenobarbital, 20 mg.; ergotamine tar- trate, 0.3 mg.; levorotatory alkaloids of bella- donne, 0.1 mg.	Sandoz Pharmaceuticals.
Do	Tablet: Phenobarbital, 40 mg.; ergotamine tar- trate, 0.6 mg.; levorotatory alkaloids of bella- donna, 0.2 mg.	Do.
Beplete with belladonna elixir	Elixir (4 cc): Phenobarbital, 16 mg.; vitamin Bi, 1.5 mg.; vitamin Ba, 1 mg.; vitamin Ba, a,33 mg.; vitamin Bi, 1.66 mg.; niacinamide, 10 mg.; pantothenol, 0.2 mg.; belladonna abalda 0.2 mg	Wyeth Laboratories.
Bexadonna	Tablet: Phenobarbital, 16 mg.; homatropine methylbromide, 10 mg.; hyoacine hydrobro- mide, 0.0065 mg.; hyoscyamine sulfate, 0.1 mg.	Bexar Pharmaceuticals.
Bilamide		Norgine Laboratories, Inc
Binitrin		The Vale Chemical Co., In
Bioxatphen		The Zemmer Co.
Bismuth, belladonna, and	Capsule: Phenobarbital, ¼ gr.; bismuth sub- gallate, 5 gr.; extract belladonna leaf, ¼ gr.	The Bernard Co.
Buffadyne A-S	Tablet: Amobarbital, 15 mg.; aspirin, 800 mg.; phenacetin, '160 mg.; caffeine, 80 mg.; hom- atropine methylbromide, 2.5 mg.; aluminum hydroxide gel, 75 mg.; magnesium hydroxide, 45 mg.	Lemmon Pharmacal Co.
Buffadyne with barbiturates	Tablet: Secobarbital sodium, 8 mg.; amobar- bital, 8 mg.; aspirin, 300 mg.; phenacetin, 150 mg.; caffeine, 30 mg.; aluminum hydroxide gel, 75 mg.; magnesium hydroxide, 45 mg.	Do.
Buncsis	Tablet: Butabarbital sodium, 10 mg.; hom- atropine methylbromide, 2.5 mg.; magnesium hydroxide, 200 mg.	McNeil Laboratories, Inc.
Buren	Tablet: Rutabarbital, 15 mg.; phenazopyridine hydrochloride, 150 mg.; scopolamine hydro- bromide, 0.0065 mg.; stropine sulfate, 0.0194 mg.; hyorcysmine sulfate, 0.1037 mg.	B. F. Ascher & Co., Inc.
Burtizem	Tablet : Butaharbital sodium, 10 mg.; reservine, 0.1 mg.; rutin, 20 mg.; mannitol hexanitrate, 30 mg.	
Butabarbital and hyoscy- amine sulfate	Tablet or elixir (5 ec.): Butabarbital, 15 mg.; hyoscyamine sulfate, 0.125 mg.	_
Do	Capule: Butabarbital, 45 mg.; hyoscyamine sulfate, 0.375 mg.	Do.
Butibel	Tablet or elixir (5 ec.): Butnbarbital sodium, 15 mg.: belladonna extract, 18 mg. (hyoscyamine sulfate, 0.128 mg.; hyoscine hydrobromide, 0.027 mg.; atropine sulfate, 0.067 mg.).	Dò.

Trade name or other designation	Compositi 👁	Manufacturer or supplier
Butibel R-A	Tablet: Hutabarbital sodium, 30 mg.; bella- donna extract, 30 mg.	Do.
Butibel-gel auspension .	Suspension (15 cc.): Butsharbital sodium, 7.5 mur.; belladonna extract, 7.5 mr. (total alka- hoids 0.187 mg.); activated attapulgite, 1.5 mg.; pectin, 75 mr.	McNeil Laboratories, Inc.
Butibel-gel tablets	Tablet: Butabarbital sodium, 7.5 mg.; bella- donna extract, 7.5 mg. (total alkaloids 0.0935 mg.); activated attapulgite, 500 mg.; pectin, 45 mg.	Do.
Butibel-Zyme		Do.
Butigetic	Tablet: Butabarbital sodium, 15 mg.; acetami- nophen, 200 mg.; phenacetin, 150 mg.; caf-	Do.
Cafergot P-B	mine tartrate, 1 mg.; caffeine, 100 mg.; levo- rotatory alkaloids of belladonna, 0.125 mg.	
Do	Suppository: Pentobarbital, 60 mg.; ergotamine tartrate, 2 mg.; caffeine, 100 mg.; levorota- tory alkaloids of belladonna, 0.25 mg.	Do.
Cal-Ma-Phen	Tablet: Phenobarbital, ¹ / ₄ gr.; calcium-carbo- nate, 5 gr.; magnesium hydroxide, 5 gr.; atropine sulfate, 1/300 gr.	Physicians Supply Co.
Cantil with phenobarbital.	Tablet: Phenobarbital, 16 mg.; mepenzolate bromide, 25 mg.	Lakeside Laboratories, Inc.
Carbonates No. 3 with phenobarbital and atro- pine	Tablet: Phenobarbital, 8 mg.; atropine sulfate, 0.11 mg.; calcium carbonate, 224 mg.; mag-	P. J. Noyes Co.
Cardalin-Phen		Mallinckrodt Chemical Works.
Cardilate-P	Tablet: Phenobarbital, 15 mg.; crythrityl tetra- nitrate, 10 mg.	Burroughs Wellcome & Co. (U.S.A.) Inc.
Cholarace	Tablet: Pentoba bital, 27.5 mg.; oxtriphylline, 200 mg.; racephedrine, 20 mg.	Warner-Chilcott Labora- tories.
Co-Elorine 25		Eli Lilly and Co.
Co-Elorine 100		Do.
Cold Preparation, special _	Tablet: Phenobarbital, 8.1 mg.; chlorphenira- mine maleate, 2 mg; pseudoephedrine hydro- chloride, 60 mg.; salicylamide, powder, 800 mg.	Knight Pharmacal Co.
Covadil		The Blue Line Chemical Co
Dactil with phenobarbital.	Tablet: Phenobarbital, 16 mg.; piperidolate hydrochloride, 50 mg.	Lakeside Laboratories, Inc.
Dainite		Mallinckrodt Chemical Works.
Dainite-Kl	Tablet: Phenobarbital, ¼ gr.; aminophylline, 3 gr.; eyhedrine hydrochloride, ¼ gr.; potas- sium iodide, 5 gr.; aluminum hydroxide gel, dried, 2½ gr.; benzocaine, ¼ gr.	Do.
Dainite Night	Tablet: Phenobarbital, % gr.; pentobarbital sosium, % gr.; aminophylline, 4 gr.; alumi- num hydroxide gel, dried, 2% gr.; benzo- caine, % gr.	
Daricon PB	Tablet: Phenobarbital, 15 mg.; oxyphencycli- mine hydrochloride, 6 mg.	Pfizer Laboratories.
Distraegus	 Tablet: Diallylbarbituric acid, ¼ gr.; nitro- glycerine, 1/250 gr.; sodium nitrite, 1 gr.; tincture crataegus, 2 minims. 	Buffington's, Inc.

Trade name or other designation	Composition	Manufacturer or supplier
Dis-Tropine	Tablet: Diallylbarbituric acid, ½ gr.: atropine suifate, 1/300 gr.: magnesium carbonate, 2½ gr.; calcium carbonate, 3½ gr.; bismuth subcarbonate, 1 gr.	Do.
	Capsule: Phenobarbital, 1/4 gr.; diphenylhy- dantoin acdium, 0.1 gm.	Parke, Davis & Co.
Do	Capsule: Phenobarbital, 1/2 gr.; diphenylhy- dantoin sodium, 0.1 gm.	Do.
Dolonil	Tablet: Butabarbital, 15 mg.; phenazopyridine hydrochloride, 150 mg.; hyoscyamine hydro- bromide, 0.3 mg.	Warner-Chilcott Laboratories.
Donabarb	Tablet: Phenokarbital, 1, gr.; powder extract belladonna, 34 gr.	Paul B. Elder Co., Inc.
Donaphen, new special donaphen	Tablet: Phenobsrbital, 15 mg.: atropine sulfate, 0.024 mg.: scopolamine hydrobromide, 0.0072 mg.: hyoscyamine hydrobromide, 0.128 mg.	Burt Krone Co.
•	Elixir (5 cc.): Phenobarbital, 16.2 mg.; hyos- cyamine hydrobromide, 0.1037 mg.; atropine sulfate, 0.0194 mg.; hysocine hydrobromide, 0.0055 mg	North American Pharmacal, Inc.
	Tablet: Phenobarbital, 8.1 mg.; phenazopyri- dine hydrochloride, 50.0 mg.; methenamine mandelate, 500 mg.; hyoscyamine sulfate, 0.0519 mg.; atropine sulfate, 0.0097 mg.; hyoscine hydrobaomide 0.0023 mg.	A. H. Robins Co., Inc.
	Indict: Phenobarbital, 15 mg.; hyoscyamine sulfate, 0.1 mg.; atropine sulfate, 0.02 mg.; scopolamine hydrohromide, 8 wg.	Lemmon Pharmacal Co.
Dormital-HM	Tablet: Phenobarbital, 3/4 gr.; homatropine methylbromide, 1/84 gr.; strontium bromide, 1 gr.	Buffington's Inc.
Dynapin with phenobar- bital	Tablet: Phenobarbital, 15 mg.; nitroglycerin, 0.5 mg.; pentaerythritol tetranitrate, 15 mg.	Key Pharmacal Co.
Elmaloin with phenobar- bital	0.5 mg.; pentaerythritol tetranitrate, 15 mg. Capsule: Phenobarbital, 15 mg.; diphenylhy- dantoin, 1½ gr.	Paul B. Elder Co., Inc.
Ephedrine and sodium phenobarbital Ephedrine sulfate and	Tablet: Sodium phenobarbital, ¼ gr.; ephed- rine sulfate, ¼ gr.	The Vale Chemical C
phenobarbital	Tablet: Phenobarbital, 15 mg.; ephedrine sul- fate, 25 mg.)	The Zemmer Co.
Ephedrine with phenobar- bital	Tablet: Phenobarbital, ¼ gr.; ephedrine sul- fate, ¾ gr.	P. J. Noyes Co.
Ercafital	fate, % gr. Tablet: Phenobarbital, 7.5 mg.; ergotamine tartrate, 0.5 mg.; caffeine, 60 mg.	The Blue Line Chemic Co.
Ethrava-trate	Tablet: Mepholarbital, 10 mg.; pentaerythrityl tetranitrate, 20 mg.; ethaverine, hydrochlo- ride, 30 mg.	North American Pharmacal, Inc.
Eu-Phed-Amin	Tablet: Phenobarbital, 30 mg.; aminophylline, 0.1 gm.; ephedrine sulfate, 30 mg.; extract	Warren-Teed Pharmace ticals Inc.
	Tablet: Phenobarbital sodium, 30 mg.; ephed- rine sulfate, 30 mg.; extract euphorbis, 0.12 gm.	Warren-Teed Pharmace ticals Inc.
Fensobel	Tablet: Phenobarbital, 8.1 mg.; belladona ex- tract, 2.95 mg.; aluminum hydrochloride gel, dried, 63 mg.; magnesium trisilicate. 63 mg.; bismuth subcarbonate, 32.5 mg.; magnesium carbonate, 252 mg.; precipitated calcium car- bonate, 203.5 mg.; malt diastase, 12.5 mg.; peppermint oil, 3 mg.	United States Vitamin Pharmaceutical Corp.
Franci	Tablet: Phenobarbital, 8 mg.; theophylline, 130 mg.; benzylephedrine hydrochloride, 32 mg.	Winthrop Laboratories.
Homechol	Tablet: Pentobarbital sodium, 8.0 mg.; homa- tropine methylbromide, 2.5 mg.; dehydrocho- lic acid, 60.0 mg.; ox bile extract, 150.0 mg.	Lemmon Pharmacal Co.
Homopent	Tablet: Pentobarbital sodium, 15 mg.; homat- ropine methylbromide, 2.5 mg; magnesium trialicate. 300 mg.	Lemmon Pharmacal Co.
H-P-A (modified)	Tablet: Phenobarbital, ¼ gr.; aspirin, 5 gr.; extract hyoscyamus, ¼ gr.	Paine Drug Co.

Trade name or other designation	Composition	Manufacturer or supplies
Hybephen	Tablet: Phenobarbital, 15 mg.; hyoscyamine sulfate, 0.1277 mg.; atropine sulfate, 0.0233	The S. E. Massengil Co.
Hybephen elixir	amine sulfate, 0.1277 mg.; atropine sulfate	Do.
Tydrochol plus	0.0233 mg.; hyoscine hydrobromide, 0.0094 mg. Tablet: Amobarbital, 15 mg.; dehydrocholic acid, 200 mg.; scopolamine metbylnitrate,	Paul B. Elder Co., Inc.
Hytrona antispasmodic elixir	0.8 mg.; ox bile desiccated, 60 mg. Elixir (5 cc.): Phenobarbital, 16 mg.; bella- donna alkaloids, 0.2 mg.	Pitman-Moore.
lytrona antispasmodic tablets	Tablet: Phenobarbital, 16 mg.; belladonna. alkaloids, 0.2 mg.	Do.
•	Tablet: Methobarbital, 30 mg.; methscopola- mine nitrate, 2.5 mg.; d-calcium pantothen- ate, 25 mg.	Warren-Teed Pharmaceuticals Inc.
	. Tablet: Phenobarbital, 15 mg; isosorbide dini- trate, 10 mg.	Ives Laboratories, Inc.
sufranol	 Tablet: Phenobarbital, 8 mg.; theophylline, 130 mg.; benzylephedrine, 32 mg.; isoprotere- nol hydrochloride. 10 mg. 	Winthrop Laboratories.
sufranol, mild	Tablet: Phenobarbital, 8 mg.; theophylline, 130 mg.; benzylephedrine, 32 mg.; isoproterenol hydrochloride, 5 mg.	Do.
suprel compound elixir	Elixir (15 cc.): Phenobarbital, 6 mg.; isopro- terenol hydrochloride, 2.5 mg.; ephedrine sulfate, 12 mg.; theophylline, 45 mg.; potas- sum india.	Da.
aphebel	Tablet : Phenobarbital, ½ gr. ; belladonna root, ½ gr. ; kaolin colluidal, 7½ gr.	Paul B. Elder Co., Inc.
anumodic	 Tablet: Phenobarbital, 1/2 gr.; belladonna root, ¹/₂ gr.; kaolin colloidal, 7/2 gr. Tablet: Phenobarbital, 8 mg.; methscopola- mine nitrate, 2 mg.; cellulase, 9 mg.; pan- crestin, 500 mg.; glutamic acid hydrochlo- ride, 200 mg.; ox bile extract, 100 mg.; pepsin, 150 mg. 	Dorsey Laboratories.
avatrate	Tablet: Phenobarbital sodium, ¼ gr.; vera- trum veride, ¼ gr.; mistletoe, ½ gr.; hawthorn tincture, 30 minims; sodium ni- tuities and the sodium ni- tuit	Key Pharmacal Co.
	Tablet: Phenoparbital, 16 mg.; potassium	Laser Inc.
iophyllin	fabict: Phenobarbital, 15 mg.; aminophyllin,	G. D. Searle & Co.
brax	Capsule: Chlordiazepoxide hydrochloride 5	Roche Laboratories.
	theophylline, 50 mg.; ephedrine hydrochlo-	Mallinckrodt Chemical Works.
uftodii tablets	Tablet : Phenobarbital, 16 mg.; theophylline, 100 mg.; ephedrine hydrochloride, 24 mg.; glyceryl gualacolate, 200 mg.	Do.
ufyllin-EP	Tablet: Phenobarbital, 16 mg.; lufyllin (dy- phylline), 100 mg.; ephedrine hydrochloride; 10 mg.	Do.
agnesium hydroxide-phe- nobarbital compound	Tablet: Phenobarbital sodium, 15 mg.; mag- nesium hydroxide, 300 mg.; atroping sulfate	McNeil Laboratories, Inc.
alglyn compound	Tablet or suspension (5 cc.): Phenobarbital, 16.2 mg.; belladonna alkaloida, 0.162 mg.	Brayten Pharmaceutical Co.
anniphen	Tablet : Phenobarbital, 16 mg.; mannitol bexa- nitrate 32 mg	The Vale Chemical Co.,
	- Tablet : Phenobarbital, 16 mg.; mannitol hexa- nitrate, 32 mg.; rutin, 20 mg.	Inc. Do.
phenoparoital	Tablet: Phenobarbital, 3/4 gr.; mannitol hexa-	
Do	Tablet: Phenobarbital, ½ gr.; mannitol hexa- nitrate, ½ gr.	The Blue Line Co.
axitol	Tablet: Phenobarbital, 15 mg.; mannitol hexa- nitrate, 15 mg.; rutin, 15 mg.; ascorbic acid, 15 mg.	Burt Krone Co.

Trade name or other designation	Composition	Manufacturer or supplier
Menrium 5-2	Tablet: Chlordiazepoxide, 5 mg. and water- soluble esterfied estrogens, 0.2 mg.	Do.
Menrium 5-4	Tablet : Chlordiazepoxide, 5 mg. and water- soluble esterfied estrogens, 0.4 mg.	Do.
Menrium 10-4	Tablet: Chlordiazepoxide, 10 mg. and water- soluble esterned estrogens, 0.4 mg.	Do.
Meprane phenobarbital	Tablet: Phenobarbital, 16 mg.; promethestrol dipropionate, 1 mg.	
Mesopin-PB	Tablet or clixir (5 cc.): Phenobarbital, 15 mg.; homatropine methylbromide, 5 mg.	
Metamine with butabar- bital	Tablet: Butabarbital, 16.2 mg.; trolnitrate	Pfizer Laboratories.
Do	Tablet: Butabarbital, 48.6 mg.; trolnitrate phosphate, 10 mg.	Do.
Mexal	Tablet : l'henobarbital, 16 mg. ; mannitol hexa- nitrate, 32 mg.	The S. E. Massengill Co.
Milprem-200		Wallace Pharmaceuticals.
Milprem-100		Do.
Milpath-200	Tablet: Meprobamate, 200 mg.; tridihexethyl chloride, 25 mg.	Do.
Milpath-100	Tablet: Meprobamate, 400 mg.; tridihexethyl chloride, 25 mg.	Wallace Pharmaceuticals.
Miltrate-10	Tablet: Meprobamate, 200 mg.; pentaerythritol tetranitrate, 10 mg.	Do.
Miltrate-20	Tablet: Meprobamate, 200 mg.; pentaerythritol tetranitrate, 20 mg.	Do.
Monomeb	bromide 5 mg	
Mudrane	Tablet: Phenobarbital, 21 mg.; potassium iodide, 195 mg.; aminophylline, 130 mg.; ephedrine hydrochloride, 16 mg.	Wm. P. Poythress & Co., Inc.
Mudrane GG elisir	Elixir (5 cc.): Phenobarbital, 5.4 mg.; theo- phylline 20 mg.; ephedrine hydrochloride, 4 mg.; glyceryl guaiacolate, 26 mg.	Do.
Nactisol		McNeil Laboratories, Inc.
	Tablet: Phenobarbital, 15 mg.; extract haw- thorn berries, 30 mg.; extract mistletoe, 15 mg.; sodium nitrite, 60 mg.; sodium blcar-	
	bonate, 0.2 gm. Tablet: Phenobarbital, 8 mg.; dehydrocholic acid, 250 mg.; bile extract, 15 mg.; hom- atroping methylbromide, 1.2 mg.	
Nergestic	atropine methylbromide, 1.2 mg. Tablet: Phenobarbital, 8 mg.; atröpine sul- fate, 0.10 mg.; magnesium trisilicate, 0.5 gm.	The S. E. Massengill Co.
Nitrased	Tablet: Secobarbital, 15 mg.; nitroglycerin, 0.4 mg.; pentaerythrityl tetranitrate, 15 mg.	Lemmon Pharmacal Co.
Nophesan tablets	Tablet: Phenobarbital, 8 mg.; acetylsalicylic acid, 300 mg.	P. J. Noyes Co.
Novalene	Tablet: Phenobarbital, 16 mg.; ephedrine sul- fate, 24 mg.; potassium iodide, 162 mg.; calcium lactate, 162 mg.	Lemmon Pharmacal Co.
	extract, 7.5 mg;; dehydrochloric acid, 32 mg;; desoxycholic acid, 32 mg;; ox bile ex- tract, 65 mg;; sorbitan mono-oleate, 160 mg;; oleic acid, 180 mg.	
Paminal elixir	Elixir (5 cc.): Phenobarbital, 8 mg.; methaco- polamine bromide, 1.25 mg.	The Upjohn Co.
Pamine PB elixir	Elixir (5 cc.): Phenobarbital, 8 mg.; methsco- polamine bromide, 1.25 mg.	Do.
Pamine PB, half strength_	Tablet: Phenobarbital, 8 mg.; methscopola- mine bromide, 1.25 mg.	Do.
Pediatric piptal anti- pyretic	Solution (0.6 cc.): Phenobarbital, 3 mg.; pipenzolate bromide, 5 mg.; acetaminophen, 60 mg.	Lakeside Laboratories, Inc.

Trade name or other designation	Composition	Manufacturer or supplier
Pediatric piptal with phenobarbital	Solution (0.5 cc.): Phenobarbital, 8 mg.; pipensolate bromide, 2 mg.	Do.
Pencetylon	Tablet: Phenobarbital, ¼ gr.; acetylsalicylic acid, 5 gr.	Paul B. Elder Co., Inc.
Pentaerythrityl tetranitrate with phenobarbital	Tablet: Phenobarbital, 16 mg.; pentaerythrityl tetranitrate, 10 mg.	
Do	Tablet: Phenobarbital, 16 mg.; pentaerythrityl tetranitrate, 20 mg.	Do.
Pentratrol with pheno- barbital	Tablet: Phenobarbital, 15 mg.; pentaerythrityl tetranitrate, 10 mg.	North American Phar- macal Co.
Pentraline	Tablet: Butabarbital sodium, 10 mg.; reser- pine, 0.05 mg.; pentaerythrityl tetranitrate, 10 mg.	McNeil Laboratories, Inc.
Perbusem	Tablet: Butabarbital sodium, 15 mg.; pentaery- thrityl tetranitrate, 10 mg.	The Zemmer Co.
Peribar L-A No. 1	Tablet: Phenobarbital, 48.6 mg.; pentaery- thrityl tetranitrate, 30 mg.	Whittier Laboratories, Inc.
Peritrate with pheno- barbital	Tablet: Phenobarbital, 15 mg.; pentaerythrityl tetranitrate, 10 mg.	
Do		Do.
Peritrate with pheno- barbital SA	Tablet: Phenobarbital, 45 mg.; pentaerythrityl tetranitrate, 80 mg.	Do.
Phedorine		Buffington's Inc.
Phenobarbital and atropine	Tablet: Phenobarbital, ¼ gr.; atropine sulfate, 1/500 gr.	Co.
Do	do	Meyers & Co. Paine Drug Co.
Do Do	Tablet: Phenobarbital, ¼ gr.; atropine sulfate, 1/250 gr.	
Phenobarbital with atro- pine sulfate	Tablet: Phenobarbital, 8 mg. ; atropine sulfate, 0.06 mg.	
Phenobarbital with atro- pine sulfate No. 2	Tablet: Phenobarbital, 15 mg.; atropine sul- fate, 0.12 mg.	The Zemmer Co.
Phenobarbital and atro- pine sulfate	Tablet: Phenobanbital, ¼ gr.; atropine sulfate, 1/200 gr.	Buffington's Inc.
Phenobarbital and atro- pine No. 1	Tablet: Phenobarbital, 16 mg. ; atropine sulfate, 0.13 mg.	Pitman-Moore.
Phenobarbital and atro- pine No. 2	Tablet: Phenobarbital, 8 mg.: atropine sulfate,	Do.
Phenobarbital and atro- pine tablets	Tablet: Phenobarbital, 8 mg.; atropine sulfate,	P. J. Noyes Co.
Do		Do.
Phenobarbital and atro- pine tablets No. 2	Tablet : Phenobarbital, 1/2 gr. ; atropine sulfate,	Do.
Phenobarbital and atro- pine tablets No. 8	Tablet: Phenobarbital, ½ gr.; atropine sulfate,	
Phenobarbital and bella- donna	Tablet: Phenobarbital, ¼ gr.; belladonna leaves	The Vale Chemical Co., In
Do	. Tablet: Phenobarbital, ¼ gr.; belladonna ex-	raine Drug Co.
Do	tract, ¾ gr. Tablet: Phenobarbital, 16 mg.; belladonna ex- tract, 8 mg.	Eli Lilly and Co.
Phenobarbital and bella- donna No. 2	Tablet: Phenobarbital, 34 gr.; belladonna ex- tract, 3% gr. (alkaloids 0.00156 gr.)	
Phenobarbital with man- nitol hexanitrate	Tablet: Phenobarbital, 7.5 mg., mannitol hexa- nitrate, 15 mg.; ascorbic acid powder, 25 mg.; rutin, 25 mg.	Paul B. Elder Co., Inc. (Harold M. Harter, D. V. M.)
Phenobarbital with man- nitol hexanitrate	Tablet: Phenobarbital, 34 gr.; mannitol nexa- nitrate, 34 gr.	 Meyer Drug & Surgical Supply Co.
Phenobarbital sodium atro pine No. 1	 Tablet: Phenobarbital sodium, 8 mg.; atropine solfate, 60 ug. 	_
Phenobarbital sodium atro pine No. 2		- 4/0.

Trade name or other designation	Composition	Manufacturer or supplier
Phenobarbital sodium atro-	Tablet: Phenobarbital sodium, 20 mg.; atropine sulfate, 200 ug.	Do.
pine No. 3 Phenobarbital and sodium nitrite	Tablet: Phenobarbital, ¼ gr.; sodium nitrite, 1 gr.	P. J. Noyes Co.
Phenobarbital theocalcin	Tablet: Phenobarbital, 15 mg.; theobromine	Knoll Pharmaceutical Co.
Phenodonna tablets	calcium salicylate, 0.5 gm. Tablet: Phenobarbital, ½ gr.; tincture bella- dunma 6 minime	Flint Medical & Surgical Supply Co.
Phenodrox	donna, 6 minima. Tablet: Phenobarbital, 14 gr.; atropine sulfate, 1/500 gr.; magnesium trisilicate, 4 gr.; alu- minum hydroxide gel, dried, 4 gr.	North American Pharmaca Inc.
Phyldrox	Tablet: Phenobarbital, 15 mg.; neothylline, 100 mg.; ephedrine sulfate, 25 mg.	Lemmon Pharmacal Co.
Piptal PHB elixir	Elixir (5cc.): Phenobarbital, 16 mg.; pipenzo- late bromide, 5 mg.	Lakeside Laboratories, Inc.
Piptal PHB tablets	Tablet: Phenobarbital, 16 mg.; pipenzolate bro- mide, 5 mg.	Do.
Prantal with phenobarbital	Tablet: Phenoharbital, 16 mg.; diphemanil methylsulfate, 100 mg.	
Premarin with pheno- barbital	Tablet: Phenobarbital 32 mg.; conjugated estro- gens-equine, 6.626 mg.	Ayerst Laboratories.
Probanthine with pheno- barbital	Tablet: Phenobarbital, 15 mg.; probanthine, 15 mg.	G. D. Searle & Co.
Probital	Tablet: Phenobarbital, 15 mg.; probanthine, 7.5 mg.	Do.
Propenite		The Zemmer Co.
Prydonnal Spansule	Capsule: Phenobarbital, 65 mg.; belladonna al- kaloids, 0.4 mg. (hyoscyamine sulfate, 0.305 mg.; atropine sulfate, 0.06 mg.; scopolamine hydrobromide, 0.035 mg.).	Smith Kline & French Laboratories.
Quadrinal	Tablet: Phen Sharbital, 24 ng.; ephedvine hy- drochloride, 24 mg; theophylline calcium sal- icylate, 130 mg.; patassium iodide, 300 mg.	Knoll Pharmaceutical Co.
Do	Suspension (5 cc.): Phenobarbital, 12 mg.; cph. Jrine hydrochloride, 12 mg.; theophylline calcium salicylate, 65 mg.; potassium iodide, 160 mg.	Do.
Quintrate with nitro- glycerin and phenobarbital Quintrate with pheno- barbital Do	Tablet: Phenobaltital, 15 mg.; pentacrythrityl fetranitrate, 20 mg.; nitroglycerine, 0.4 mg. Tablet: Phenobarbital, 15 mg.; pentacrythrityl tetranitrate, 10,mg. Tablet: Phenobarbital, 15 mg.; pentacrythrityl	(Giynn A. Beard).
Robinul-PH	tetranitrate, 20 mg. Tablet : Phenobarbital, 16.2 mg.; glycopyrro-	A. H. Robins Co., Inc.
Robinul-PH forte	late, 1.0 mg. Tablet: Phenobarbital, 16.2 mg.; glycopyrro-	Do.
Ruhexatal	late, 2.0 mg. Tablet: Phenobarbital, 15 mg.; mannitol hexani- trate, 30 mg.; ascorbic acid, 10 mg; rutin, 20 mg.	Lemmon Pharmacal Co.
Rutol	Tablet : Phencharbital, \$ 0 mg.; mannitol hexa- nitrate, 16 mg.; rutin, 10 mg.	Pitman-Moore.
Salisii with phenobarbital	Tablet: Phenobarhital, ¹ / ₂ gr.; acetylsalicylic acid, 5 gr.; nugnosium trisilicate, 2 gr.	Paul B. Elder Co., Inc.
Selbella	Tablet : Phenobarbital, 1; gr.; aluminum hy-	Wyeth Laboratories.
Sed-Tens	dvoxide, 5 gr.; belladonna extract, ½ gr. Tablet (12 hr.): Amoba(bital, 50 mg.; homatro- pine methylluomide, 7.5 mg.	Lemmon Phaemacal Co.
Sibena	Tablet: Butabarbital sodium, 16 mg.; simethi- cone, 25 mg.; belladonna extract, 16 mg.	Plough Laboratories, Inc.
Sodium nitrite with pheno- barbital	(total alkaloids, 0.20 mg.). Tablet: Phenobarbital sodium, ¼ gr.; sodium nitrite, 1 gr.; sodium bicarbonate, 2 gr.;	Paine Drug Co.
Do	hawthorn berries, fluid extract, ¼ minim. Tablet: Phenobarbital, ½ gr.; sodium nitrite,	Buffalo Pharmaceutical
Spasticol PB	1 gr. Tablet: Phenobarbital, 15 mg.; homatropine methylbromide, 2.5 mg.	Supply Corp. Key Pharmaceuticals, Inc.

Trade name or other designation	Composition	Manufacturer or supplier
Spastored	Tablet: Phenobarbital, 8 mg.; atropine sulfate, 0.13 mg.; calcium carbonate, 227 mg.; mag-	North American Pharmacal, Inc.
Synirin	nesium hydroxide, 162 mg. Tablet: Pentobarbital, 8 mg.; aspirin, 324 mg.	Wm. P. Poythress & Co., Inc.
rcs	Tablet : Phenobarbital, 16 mg. ; theobromine sal-	Do.
Fedral-25	icylate, 0.4 gm.; calcium salicylate, 0.06 gm. Tablet: Butabarbital, 25 mg.; theophylline, 130	Warner-Chilcott Laboratories.
Tedral S.A.	mg.; ephedrine hydrochloride, 24 mg. Tablet: Phenobarbital, 25 mg.; theophylline, 180 mg.; ephedrine hydrochloride, 48 mg.	Warner-Chilcott Laboratories.
Fensodin	Tablet : Phenobarbital, 16 mg. ; ethaverine hy- drochloride, 30 mg. ; theophylline calcium sal-	Knoll Pharmaceutical Co.
Tensophen	icylate, 200 mg. Tablet: Phenobarbital, 16 mg.; nitroglycerin, 0.26 mg.; sodium nitrite, 32 mg.; podophyl- lin, 1 mg.; extract beef bile, 16 mg.	P. J. Noyes Co.
Thedrizem	Tablet : Phenobarbital, 8 mg. ; theophylline, hy- drous, 100 mg. ; ephedrine hydrocloride, 25 mg.	The Zemmer Co.
Theobarb	- Tablet: Phenobarbital, 32 mg.; theobromine, 325 mg.	Mallinckrodt Chemical Works.
Theobarb-R	- Tablet: Phenobarbital, 10 mg.; reserpine, 0.1 mg.; theobromine, 324 mg.	Ъо.
Theobarb special	- Tablet: Phenobarbital, 16 mg.; theobromine, 325 mg.	Do.
Theobromine and pheno- barbital	Tablet : Phenobarbital, 16 mg.; theobromine, 0.3 gm.	P. J. Noyes Co.
Theobromine-phenobarbital	Tablet: Phenobarbital, 30 mg.; theobromine, 0.3 gm.	The S. E. Massengill Co.
Do		The Upjohn Co.
Theobromine-phenobarbital compound	Tablet : Phenobarbital, 1/4 gr. ; theobromine, 21/4	Do.
Theobromine with pheno- barbital No. 1	Tablet: Phenobarbital, 15 mg.; theobromine,	Buffington's, Inc.
Theobromine and sodium acetate with phenobar- bital	Tablet : Phenobarbital, ¼ gr. ; theobromine and sodium acetate, 3 gr.	Paul B. Elder Co., Inc.
Theobromine sodium saliciylate with pheno-	Tablet: Phenobarbital, 15 mg.; theobromine so- dium salicylate, 300 mg.	The Zemmer Co.
barbital Theocardone No. 1	. Tablet : Phenobarbital, 15 mg. ; theobromine, 800 mg.	Haack Laboratories, Inc.
Theocardone No. 2		Haack Laboratories, Inc.
Theodide		The Vale Chemical Co., In
Theoglycinate with pheno- barbital	Tablet: Phenobarbital, 16 mg.; theophylline-	Brayten Pharmaceutical C
Theoglycinate with race- phedrine and phenobar-	Tablet: Phenobarbital, 16 mg.; theophylline- sodium glycinate, 324 mg.; racephedvine hy-	Brayten Pharmaceutical (
bital Theoplaphen	 Tablet: Phenobarbital, 15 mg.; theohromine sodium salicylate, 0.2 gm.; calcium lactate. 	The S. E. Massengill Co.
Theominal	0.1 mg. Tablet: Phenobarbital, 32 mg.; theobromine,	Winthrop Laboratories.
Theominal M	320 mg. . Tablet: Phenoharbital, 16 mg.; theobromino, 320 mg.	Do.
Theominal R S		Do.
Theophen	 Tablet: Phenobarbital, ¼ gr.; theobromine so- dium salicylate, 5 gr.; calcium carbonate, 	The Vale Chemical Co., In
Theorate	2½ gr. . Tablet: Phenobarbital, 16.2 mg.; theobromine, .324 mg.	Whittier Laboratories, In
Thymodyne -	324 mg. Tablet : Phenobarbithi, 32 mg. ; theophylline an- hydrous, 130 mg.; ephedrine sulfate, 24 mg.	P. J. Noyes Co.
Trocinate with pheno- barbital	Twhet: Phen barbital, 16 mg.; thiphenamil hydrochloride, 100 mg.	Wm. P. Poythress & Co., Inc.

Trade name or other designation	Composition	Manufacturer or supplier
Tricoloid	Tablet: Phenobarbital, 16 mg.; tricyclamol	Burroughs Wellcome & Co.
Triop hen	chloride, 50 mg. Tablet: Phenobarbital, ½ gr.: stropine sulfate,	The Vale Chemical Co., Inc.
Valpin-PB	1/500 gr.; magnesium trisilicate, 7 gr. Tablet or elixir (5 cc.): Phenobarbital, 8 mg.;	Endo Laboratories, Inc.
Vasorutin	anisotropine methylbromide, 10 mg. Tablet: Diallylbarbituric acid, ¼ gr.; nitro- glycerine, 1/250 gr.; sodium nitrite, 1 gr.;	Buffington's, Inc.
Veralzem	tincture crataegus, 2 minims; rutin, 20 mg. Tablet: Phenobarbital, 15 mg.; veratrum viride,	The Zemmer Co.
Veratrite	50 mg.; sodium nitrite, 60 mg. Tablet: Phenobarbital, ¼ gr.; cryptenamine, 40 CSR (carotid sinus reflex) units; sodium	
Veritag	nituite, 1 gr. Tablet: Phenobarbital, 16 mg.; veratrum viride,	S. J. Tutag and Co.
Vertegus	 a) OSA (tabtin into into inter, units, south inter, 1 gr. Tablet: Phenobarbital, 16 mg.; veratrum viride, 40 mg.; sodium nitrite, 65 mg. Tablet: Phenobarbital, 1/4 gr.; weratrum viride, 1/4 gr.; sodium nitrite, 1 gr.; mistletce, 1/4 gr.; tablet: Phenobarbital, 1/5 mg.; rutin, 20 mg.; 	Burt F.rone Co.
Veruphen	gr.; hawthorn berries, ½ gr. Tablet: Phenobarbital, 15 mg.; rutin, 20 mg.; veratrum viride, 15 mg.; sodium nitrite, 60 mg.	The Zemmer Co.
Viritin	. Tablet: Phenobarbital, 15 mg.; mannitol hexa- nitrate, 30 mg.; veratrum viride alkaloids,	Lemmon Pharmacal Co.
W-T	 5 mg²; rutin, 20 mg. Powder (4 gm.): Phenobarbital, 15 mg.; bella- donna extract, 10 mg. (0.12 mg. belladonna alkaluids): benzocaine, 15 mg.; calcium car- bunate, 1.55 gm.; magnesium oxide, 0.5 gm.; 	CMIR, INC.
W-T	aluminum hydroxide gal dried 60 mg.	Do.
Xaniophen	chlorophyll extract, 1%. Tablet: Phenobarbital, 16.2 mg.; theobromine, 162 mg.; ethylenediamine dihydriodide, 32.4	Pitman-Moore.
Zallogen compound	mg. - Tablet: Phenolarbital, 8 mg.; tocamphyl, 75	The S. E. Massengill Co.
Zantrate	ng.; homatropine methylbromide, 2.5 mg. Tablet: Cyclogentenylallybarbituric acid, 1/2 gr.; ephedrine sulfate, 3/2 gr.; theophylline	The Upjohn Co.
Zem-Dab	anhydrous, 2 gr. Tablet: Butabarbital sodium, 10 mg.; debydro- chulic acid, 60 mg.; ox bile desiccated, 120	The Zemmer Co.
No. 23	mg.; homatropine methylbromide, 2.5 mg. Tablet: Phenobarbital, ½ gr.; aminophylline,	Stayner Corp.
No. 35	3 gr. Tablet: Phenobarbital, ¼ gr.; aminophylline,	Do.
No. 36	 gr.; ephedrine sulfate, 3g gr. Tablet: Pentabarbital sodium, 4g gr.; ephedrine sulfate, 3g gr.; aminophylline, 3 gr. 	Do.
No 65	. Tablet: Phenoharbital, 15 gr.: extract pena-	· 1/0.
No. 66	donna, ¼ gr. _ Tablet: Phenobarbital, ¼ gr.; extract bella-	Do.
No. 75	donna, ¼ gr. - Tablet : Phenobarbital, ¼ gr. : belladonna, ¼ gr.	Bariatric Corp.
No. 88	 Tablet: Phenobarbital, ¼ gr.; aminophylline, 1.5 gr. 	, Stayner Corp.
No. 89	Tablet: Phenobarbital, ½ gr.; aminophylline,	
No. 111	_ Tablet: Phenobarbital, 14 gr.; ephedrine sul-	- Do.
No. 186	fale, % gr. - Tablet: Phenobarbital, 20 mg.; homatropine methylbumide 5 mg	e Do.
No. 643	methylbromide, 5 mg. Tublet : Phenobarbital, ½ gr.; theophylline, 2 gr.; ephedrine hydrochloride, 3g gr.	Do.
Rx. No. 4104	Tablet: Phenotherbital, '4 gr.; calcium carbon. atc, 73' gr.; magnesium oxide, 4 gr.; atro pine sulfate, 1/300 gr.	. The Zemmer Co.
Rx. No. 4105	 Tablet: Phenobabical, % gr.; calcium carbon ate, 10 gr.; attopine sulfate, 1/300 gr. 	• Do.

Trade name or other designation	Composition	Manufacturer or supplier
Rx. No. 4108	Capsule: Phenolarbital, ¼ gr.; atropine sul- fate, 1/300 gr.; calcium carbonate, 6½ gr.; magnesium oxide, heavy, 2 gr.	Do.
Rx. No. 4123	Capsule: Phenobarbital, ¼ gr.; bismuth sub- galate, 6 gr.; extract belladonna, ¼ gr.	Do.
Rx. No. 4126	Capsule: Pentobarbital sodium, 15 mg.; extract belladonna, 10 mg.	Do.
Rx. No. 4143	Capsule: Phenobarbital, 14 gr.; aminophylline, 1.5 gr.; potassium iodide, 1 gr.	Do.
Rx. No. 4152	Tablet: Phenobarbital, ½ gr.: atropine sulfate, 1/200 gr.	Do.
Rx. No. 4155	Tablet: Phenobarbital, ½ gr.; atropine sulfate, 1/1000 gr.; aluminum hydroxide gel, 3% gr.; kaolin, 3% gr.	Do.
Rx. No. 4170	Tablet: Phenobarbital. 1/2 gr.; stropine sulfate, 1/200 gr.; calcium carbonate, 10 gr.	Do.
Rx. Nó. 4184	Capsule: Sodium butabarbital, 15 mg.; bella- donna extract, 15 mg.	Do.

EXCEPTED PRESCRIPTION DRUGS-Continued

(Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.22; filed Jul 9, 1974, 9:29 am: Unpublished; filed Jun 9, 1977, 8:55 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-9 Application for exclusion of stimulant or depression compound; revocation

Authority: IC 35-48-3-1 Affected: IC 35-48-2-1

Sec. 9. Application for exclusion of a stimulant or depressant compound. (a) Any person seeking to have any non-narcotic substance which may, under the Federal Food, Drug, and Cosmetic Act or state law, be lawfully sold over-the-counter without a prescription, excluded from any schedule, pursuant to IC 1971, 35-24.1-2-1(g) [*Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.]* as amended, may apply to the Indiana Board of Pharmacy for such exclusion.

(b) An application for an exclusion under this section shall be approved by the Board, provided that such application shall contain a copy of the Administration's approval for such request for an exclusion from the Federal Controlled Substances Act.

(c) Within a reasonable period of time after the receipt of an application for an exclusion under this section, the Indiana State Board of Pharmacy shall notify the applicant of its acceptance or non-acceptance of the application, and if not accepted, the reason therefor. The Indiana State Board of Pharmacy need not accept an application for filing if any of the requirements prescribed in paragraph (b) of this section is lacking or is not set forth so as to be readily understood. If the applicant desires, he may amend the application to meet the requirements of paragraph (b) of this section.

(d) The Indiana State Board of Pharmacy may at any time revoke any exclusion granted pursuant to IC 1971, 35-24.1-2-1(g) [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.], as amended, upon a finding that such exclusion from the Federal Controlled Substances Act was terminated, suspended or revoked. The revocation of the exclusion granted under this Act [IC 35-48] shall become effective upon the board's notifying the person to whom such exclusion was granted by certified mail of such revocation. (Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.23; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-10 Excluded nonnarcotic substances, stimulant or depressant compounds

Authority: IC 35-48-3-1 Affected: IC 35-48-2-1

Sec. 10. Excluded non-narcotic substances, stimulant, or depressant compounds. (a) The Indiana Board of Pharmacy may exclude any non-narcotic substance from a schedule if such substance may, under the Federal Food, Drug and Cosmetic Act or state law, be lawfully sold over-the-counter without a prescription pursuant to IC 1971, 35-24.1-2-1(g) [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended, the drugs which were excluded by the Bureau or Administration on January 1, 1974 under section 201(g)(1) of the Federal Controlled Substances Act (21 U.S.C. 811(g) (1)) have been excluded by the Indiana State Board of Pharmacy from the schedules of IC 1971, 35-24.1-2-4, 6, 8, 10, and 12 [Repealed by

Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.J, as amended. The exclusion of these drugs by the Indiana State Board of Pharmacy should not be construed as an adoption or rejection of the criteria by which these drugs were originally excluded. Any deviation from the quantitative composition of any of the listed drugs shall require a petition for exclusion in order for that drug to be excluded. The following is a list of the presently excluded non-narcotic substances under these regulations.

Trade name or other designation	Composition	Manufacturer or supplier
Amodrine	Tablet: Phenobarbital, 8 mg.; aminophylline, 100 mg.; racephedrine hydrochloride, 25 mg.	G. D. Searle & Co.
Bronkolixir	Elixir (5 cc.): Phenobarbital, 4 mg.; ephedrine sulfate, 12 mg.; glyceryl guaiacolate, 50 mg.; theophylline, 15 mg.	Breon Laboratories, Inc.
Bronkotabs	Tablet: Phenobarbital, 8 mg.; ephedrine sul- fate, 24 mg.; glyceryl guaiacolate, 100 mg.; theophylline, 100 mg.	Do.
Primatene	Tablet : Phenobarbital, 1/8 gr. ; ephedrine, % gr.	Whitehall Laboratories,
Rynal	Solution for Spray: dl-Desoxyephedrine HCL 0.22 ['] / ₄ ; antipyrine 0.28 ['] / ₆ ; pyrilamine maleate 0.01 ['] / ₆ : methyl dodecylbenzyltrimethyl am- monium chloride 0.02 ['] / ₄ ; glycerine dehydrate 1.50 ['] / ₆ .	Blaine Co.
Tedral	Tablet: Phenobarbital. 8 mg.; theophylline, 130 mg.; ephedrine hydrochloride, 24 mg.	Warner-Chilcott Laboratories,
Tedral Anti-H	Tablet: Phenobarbital, 8 mg.; chlorphenira- mine maleate, 2 mg.; theophylline, 130 mg.; ephedrine hydrochloride, 24 mg.	Do.
Tedral one-half strength	Tablet: Phenobarbital, 4 mg.; theophylline, 65 mg.; ephedrine hydrochloride, 12 mg.	Do.
Tedral Pediatric Suspension	Suspension (5 cc.): Phenobarbital, 4 mg.; ephedrine hydrochloride, 12 mg.; theophyl- line, 65 mg.	Do.
Tedral suppositories double strength	Suppository: Phenobarbital, 16 mg.; theophyl- line 260 mg.; ephedrine hydrochloride, 48 mg.	Do.
Tedral suppositories regular strength	Suppository: Phenobarbital, 8 mg.; theophyl- line, 130 mg.; ephedrine hydrochloride, 24 mg.	Do.
Verequad	Tablet: Phenobarbital, 8 mg.; theophylline cal- cium salicylate, 130 mg.; ephedrine hydro- chloride, 24 mg.; glyceryl guaiacolate, 100 mg.	Knoil Pharmaceutical Co.
Verequad	Suspension (5 &c.): Phenobarbital, 4 mg.; theo- phylline calcium salicylate, 65 mg.; ephedrine hydrochloride, 12 mg.; glyceryl guaiacolate, 50 mg.	Do.

EXCLUDED OVER-THE-COUNTER DRUGS

(Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.24; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-11 Exempt chemical preparations

Authority: IC 35-48-3-1 Affected: IC 35-48-2-1

Sec. 11. Exempt Chemical Preparations. (a) The chemical preparations and mixtures specifically listed in subparagraph (b) of this Section have been exempted by the Indiana Board of Pharmacy from the application of IC 1971, 35-24.1-3-2, 3, 6 and 8 *[Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.]* as amended, which preparation or mixture is intended for laboratory, industrial, educational or special research purposes and not for general administration to a human being or other animal. The exemption to be valid must be in strict compliance with the requirements imposed for the preparation or mixture prescribed in Part 1308, Section 1308.24 of Title 21 of the Code of Federal Regulations, effective January 1, 1973, and no exemption granted pursuant to this Section affects the criminal liability for illegal manufacture, distribution, or possession of controlled substances contained in the exempt chemical preparation. Distribution, possession and use of an exempt chemical preparation are lawful for registrants and non-registrants only as long as such distribution, possession or use is intended for laboratory, industrial or educational purposes and not for immediate or subsequent administration to a human being or other animal.

(b) The following preparations and mixtures in the form and quantity listed in the application submitted (indicated as the "date

of application") are designated as exempt che	emical preparations for the purposes set forth in this Section.

Manufacturer or supplier	Product name and supplier's catalog number	Form of product	Date of application
Abbott Laboratories	CEP Agarose Plates, NDC 0074- 9023-12, NDC 0074-9023-35.	Foil Pouch: 4½ by 4 inches, 6½ by 5½ in- ches.	Mar. 24, 1975
	Barbital-Acetate Buffer Powder with 1 g Sodium Azide, NDC 0074-7591-12.	Plastic Bag: 16.24 g per bag.	Do.
Do	Digoxin I 125 Imusay • diagnostic kit No. 7649.	Kit: 100 units	June 6, 1974
Do	HTSH RIA diagnostic kit No. 7504	Kit: 50 units	Sept. 25, 1974
Do	Tetrasorb-125 T-4 diagnostic kit No. 7775.	·· .	Aug. 21, 1972
Do	Irosorb-59 diagnostic kit No. 6764	Vial: 10 ml	Do.
Do	Quantisorb T-4N diagnostic kit No. 6719.	. 200°	Do.
Airwick Industries	Airkem Solidaire Green	Tube: 7 oz. and 14 oz.	Dec. 5, 1973
Do	Airkem Solidaire Gold		Do.
Do	Airwick Solidaire Citrus	Do	Do.
Do	Airkem Musketeer, Jr.		Do.
Do	Airwick Solid Natural		Do.
Do	Airwick Solid Floral		Do.
Do	Airwick Solid Lemon		Do.
Do	Airwick Solid Rose	Do	Do.
American Hospital Supply Corp. (Dade Division).	Buffered Thrombin (Bovine), Cata- log No. B4233-40, Eugiobulin Lysis Set	•	
Do	Fibrin Monomer Control, Cata- log Nos. B4233-30 and B4233-38.	Bottle: 1.5 ml	Feb. 16, 197

Manufacturer or supplier	Product name and supplier's catalog number	Form of product	Date of
Do	Moni-Trol I-X Chemistry Controls (Level I), Catalog Nos.:		T PA 100-
	(Level I), Catalog Nos.: B5106-1 B5106-5 B5106-5	Vial: 10 ml. Bottle: 25 ml.	ani. 40, 1979
	Moni-Trol II-X Chemistry Controls (Level II) Catalog Nos.:	A CALL AND A	•
	B5106-2 B5106-6 B5106-6	Vial: 5 ml Vial: 10 ml Bottle: 25 ml	Do.
Do	Owren's veronal buffer No. B4234-	Bottle: 15 ml	Jan. 22, 1973
Do	Phosphatase substrate No. B5312- 1 and No. B5312-5.	Bottle: 73 mg. dry powder.	Do.
	Serum reagent No. B4233-1 and No. B4233-2.		Do.
Do	.84233-10.	••••	Do.
Do	DATA-topens T 125 4. Buffered T thyroxine, catalog No. B5644-21.	Bottle: 55 ml	June 11, 1975
	DATA-topens T 125 4. Buffered I thyroxine, catalog No. B5644-25.		
Do 20. "As aver V off	DATA-topend T 125 4. Buffered I thyroxine, catalog No. B5644-29.	Bottle: 505 ml	Do.
Do	DATA-topens CT 125 4. Buffered I thyroxine, catalog No. B5644-40.	Bottle: 55 ml	Do.
•	DATA topens CT 125 4. Buffered I thyroxine, catalog No. B5644-45.	. •	Do.
a state in the second secon	DATA-topens T 125 4. Buffered I thyroxine, catalog No. B5644-35.	· · · · · · · · · · · · · · · · · · ·	Do.
American Hospital Supply Corp. (Harleco Division).	Barbital buffer B-1 No. 96772	7 dram vial.	
Do	double strength, pH 8.6, 0.075 m No. 938-34.		
Do	Barbital-sodium buffer salt, No. 11731.	Bottle: 250 ml	June 6, 1972
Do	Barbital-acid buffer sait, No. 1173	Bottle: 250 ml	Do.
Do	Buffer sait mixture Spinco B-1, pH 8.6, 0.05 ionic strength, No. 3947.	Vial: 12.12 grams per 29.5 x 80 mm. vial.	Sept. 15, 1971
Do	Buffer salt mixture Spinco B-2, pH 8.6, 0.075 ionic strength, No. 3948.	Vial: 18.18 grams per 29.5 x 80 mm. vial.	Do.
Do	Buffer for serum protein electro-	Vial: 10 dram	July 25, 197
	catalog No. 32307.	taining a powder to be reconstituted by	<i>May</i> 01, 131
American Monitor Corp	Qualify I Qualify II	Glass vial : 10 ml	Oct. 9, 197
Amersham/Searle	Amobarbital-2-C14, No. CFA-401_	Amnula: 110 mm x 13	Sant 10 197
······································		mm. or Vial: 38.40 mm, x 11 ml.	
Do Do	HPL Immunoassay Kit No. IM-63. Morphine (N-methyl-C14) Hydro- chloride No. CFA-363.	Bottle: 30 ml Do	May 18, 197 Mar. 27, 197
	Pentothal-S35 sodium salt, No. SJ-77.	13 mm. or Vial: 38.40 mm. x 11 mm.	
Do	Codeine (N-methyl-C14) Hydro- chloride No. CFA-421.	Ampule: 10 cc.	Mar. 27, 197

Product name and supplier's catalog number Date of Manufacturer or supplier Form of product application d-(side chain ³H) Amphetamine Ampoule: 5 mi _____ Sulfate Number TRK-444. Do Sept. 20, 1973 Lysergic acid di[1-14C] ethyla-mide, catalog No. C.F.A. 534. Do Ampoule: 0.6 mg. to July 2, 1974 8.1 mg. T-3 RIA Kit Catalog No. IM. 74_ Kit Do Nov. 4. 1974 Pheno [2-14C] barbital Catalog Ampoule: No. C.F.A. 537. curies. Do 50 micro- Nov. 5, 1974 .__Do ____. Do Ampoule: 250 micro-Do. curies. [15, 16(n)-3H] Etorphine, Cata-log No. T.R.K. 476. D٥ Ampoule: 250 micro- Nov. 19. 1974 curies. [2(n)-3H] Lysergic acid diethyl-amide, No. T.R.K. 461. Ampoule: 0.003 mg. to May Do 22. 1974 0.04 mg. [15, 16(n)-3H] Etorphine Cata- Ampoule: 1 millicurie_ Feb. log No. T.R.K. 476. Do 17. 1975 -)Δ¹-Tetrahydro [3', 5'-14C] Cannabinol Catalog No. C.F.A. Do 5'-14C] Ampoule: 10 and 50 Mar. (-5, 1975 microcuries. 538. d-[methylene ¹⁴C] Amphetamine Sulphate, catalog No. C.F.A. 544. Do Ampoule: 110 x 13 mm. June 11, 1975 T-4 RIA Kit, catalog No. IM 80. Do Kit containing: 50 Nov. 25, 1975 tests. T-4 RIA Kit, catalog No. IM-801. Kit containing: tests. Do 100 Do. T-4 RIA Kit, catalog No. IM 801A. Do Do Do. Amersham/Searle Corp. __ 26. 1974 Da [1(n)-3H] Morphine, No. T.R.K. 447. Ampoule: 0.002 mg. to Do. 0.015 mg. Do [1(n)---3H] Codeine, No. T.R.K. 448. Do. Do Diacetyl [1(n)-³H] morphine, No. T.R.K. 449. Ampoule: 0.003 mg. to Do. 0.012 mg. [1,7,8 (n)-3H] Dihydromorphine, No. T.R.K. 450. Ampoule: 0.0008 mg. to 0.008 mg. Do Do. Analytical Chemists, Inc. _ Sodium Barbital Buffer, Catalog Vial: 20.6 g Aug. 14. 1972 Nos. 1-5100 and 1-5200. Agarose Universal Electrophoresis Plate: 5 ml _____ Film, Catalog No. 1-1000. Do Do. Toxi-Disem A, 121, A-1; 122, A- Dise: ½ in x 0.2 mm_ May 2; 124, A-4. Analytical Systems 6, 1975 - 1 Toxi-Disens B, 125, B-1; 126, B-2; ____Do _____ 127, B-3; 128, B-4. Dó Do. Applied Sciences Laboratories, Inc. Mixture 1-opiates Vial: 1 ml Oct. 4, 1972 ----------- Mixture 2--stimulants Do Mixture 8--depressants Do Mixture 4--barbiturates Do Mixture 6--kit of representatives Do Opiates, Mixture 1 Number 01830 Vial: 10 ml Stimulants, Mixture 2 Number Do Do _____ Do. Do Do. Do Do. -----Do -----Do. Do Oct. 4, 1973 Do Do. 01831. Depressants, Mixture 3 Number ____Do _____ 01832. Do Do. Barbiturates, Mixture 4 Number ____Do Do Do. 01833. Allyliscbutylbarbituric acid, No. Vial: 1 ml Do Jan. 24. 1973 Alphenal, No. 01743 Amobarbital, No. 01744 Amphetamine HCL, No. 01745 Do ----- Do Do. Do ____ Do Do. Do. ----------Do ____Do

Manufacturer or supplier	Product name and supplier's or catalog number		Date of application
Do	Aprobarbital, No.: 01746	Da	. Dec
Do	Barbital, No. 01747	Do	Do. Do.
Do	Barbital, No. 01747 Butabarbital, No. 01748	Do	Do.
Do	Butethal No. 01749	Do	Do.
Do	Cocaine, No. 01750		Do.
Do	Codeine, No. 01751	Do	Do.
Do	Diallybarbituric acid, No. 01752	Do	Do.
Do Do	Etheniorvynol, No. 91753	Do	Do.
Do			Do.
	Glutethimide No 01756		Do.
Do	Glutethimide, No. 01756 Hexobarbital, No. 01757	Do	Do. Do.
Do	Hydrocodone Ritartrate, No. 01758	Do	Do.
* Da	Meneridine HCI. No 01750	The second se	Do.
Do	Meperidine HCL, No. 01759 Mephobarbital, No. 01760 Meprobarate, No. 01761	Do	Do.
Do	Manunhamata No 01761	10	Do.
Do	Mescaline, No. 01762	Do	Do.
Do	Methadone HCL, No. 01763	Do	Do. :
Do	Methamphetamine HCL, No. 01764	Do	Do.
Do	Methylphenidate, No. 01774	Do	Do.
Do	Morphine, No. 01765	Do	Do.
			Do.
Do	Pentobarbital, No. 01767	Do	Do.
Do	Phenasocine H Br, No. 01768		Do.
Applied Sciences Labora- tories, Inc.	Phencyclidine HCL, No. 01769	Vial: 1 ml.	Jan. 24, 1973
Do	Phenobarbital, No. 61776	The State	Do.
Do	Seconstruital, No. 01771	Do	Do.
Do	Thebaine, No. 01772	Do	Do.
Do	"I"htsmtylal, No. 61773	Ba	De
Beckman (Instruments, Inc. (Spince Division)	ASO buffer, pH 7.2	Tube: 2.7 grams	Aug. 31, 1973
Do Do	Beckman buffer B-1 Beckman buffer B-2	Packet: 12.1: gm Packet: 18.16 gm	Apr. 24, 1971 Do.
(diagnostic operations) ~	Human thyroid stimulating hor- mone kit, single label: No. 565185		Nov. 26, 1974
5 . av are	No. 566185	10 tests 25 tests 50 tests	
	No. 566187	50 tests	
and the second second	No. 566188 Human thyroid stimulating hor-	100 tests	· .
1		A STATE OF A	Do.
er i	No. 566174	10 tests	
٠. ١	No. 566173 No. 566174 No. 566175	25 tests	
•	ANU. 000110	100 tests	
Du	Triiodothyronine kit single label	Kit, containing :	Do.
	No. 566177 No. 566178 No. 566179	10 tests	./:
"	No. 566179	25, teste	7
Do	No. 566179 Triiodothyronine kit, double label:	Kit, containing :	Do.
*:	No. 566181	10 tests	1.0.
		50 tests	
_	No. 565183		
Du	No. 566183 Thyroxine kit, single label:	Kit, containing :	Do.
Du	No. 566183 Thyroxine kit, single label: No. 566166 No. 566166	Kit, containing:	Do.
· , ·	No. 566166	Kit, containing: 10 tests 25 tests	Do.
· , ·	No. 566166	Kit, containing: 10 tests 25 tests	
· , ·	No. 566166	Kit, containing: 10 tests 25 tests	Do. Do.
· , ·	No. 566166	Kit, containing: 10 tests 25 tests	
· , ·	No. 566166	Kit, containing: 10 tests 25 tests	
· , ·	No. 566166 No. 566167 Thyroxine kit, double label: No. 566169 No. 566169 No. 566171	Kit, containing: 10 Lests 25 tests 50 tests 10 Lests 10 Lests 25 tests 26 tests 27 tests 26 tests	Do.
· , ·	No. 566166 No. 566167 Thyroxine kit, double label: No. 566169 No. 566170 No. 566171 Digoxin kit, single label: No. 566171	Kit, containing: 10 Lests 25 tests 50 tests 10 Lests 10 Lests 25 tests 26 tests 27 tests 26 tests	
· , ·	No. 566166 No. 566167 Thyroxine kit, double label: No. 566169 No. 566170 No. 566171 Digoxin kit, single label: No. 566171	Kit, containing: 10 tests 25 tests 50 tests 10 tests 10 tests 25 tests 26 tests 27 tests 50 tests 26 tests 10 tests 10 tests 10 tests 10 tests 10 tests 10 tests	Do.
Do	No. 566166 No. 566167 Thyroxine kit, double label: No. 566169 No. 566169 No. 566171	Kit, containing: 10 tests 25 tests 50 tests 50 tests Kit, containing: 10 tests 25 tests 50 tests Kit, containing: 10 tests 25 tests 50 tests 26 tests 50 tests 10 tests 25 tests	Do.

Manufacturer or suppliers	Product name and supplier's	Form of production	Date of application
(Spectra Biologicals Di-			Aug. 11, -1972
Do	HepaScreen CEP plates, Nos. K- 742 and K-743.	Plate: 3.5" x 3.6"	Dov
Behring Diagnostics, American Hoechst Corp.	Immuno-teck II Agarose Plates	Foil pouch : 5.35 by 5.25 in.	
Der	IEP Buffer pH 8.2	Foil pouch: 5.35 by 5.25 in., 65 g	Do.
Biomedical Products Corp. Do	Barbiturate stock standard, 3-1303 Mayer's hematoxylin solution, 6- 1192	Bottle: 100 ml Bottle: 100 mi, 500 ml, 1,000 ml, 1 gal.	Apr. 18.0197
Bio-Rad Laboratories, Inc.	Bio-Rad electrophoresis buffer Barbital Buffer-Dry Pack	Bottle: 500 ml.	Dec. 14, 197
Do	Do	Package: 12.14 gm.	Do
Do	Electrophoresis buffer, dry-pack	Package: 6.15 gm.	Dec. 14, 197
Do Do	Do Do Electrophoresis buffer, dry-pack Resgent No. 3 Immunoelectrophoresis Barbital Buffer I, pH 8.6.	Dry-pack: 25.6 gm.	Aug. 6, 197
Du	Buffer I, pH 8.6. immunoelectrophoresis Barbital Buffer II, pH 8.6.	Dry-pack: 15.61 gm	Do.
Do	Immunoelectrophoresis Barbital Buffer III-a, pH 8.8.	Dry-pack: 15.07 gm	Do. '
Bio-Rescents &	Prochex No. 700-225	Vial: 25 ml.	·Mar. 9, 197
Do	Prochex No. 1, No. 701-025	<u>D</u> o	Do.
Do	Prochez No. 1, No. 701-025 Prochez No. 1 (Alternate Formu- la) No. 702-025. Prochez No. 2, No. 703-025 Prochez No. 3, No. 704-025 Prochez No. 3, No. 706-025 Prochez No. 5, No. 706-025 Prochez No. 6, No. 707-025 Prochez No. 7, No. 708-025 Prochez No. 9, No. 710-025 Prochez No. 9, No. 710-025 Prochez No. 10 (Alternate Formu- la) No. 712-025) Prochez No. 10 (Alternate Formu- la) No. 712-025)	Do	Do.
Do	Prochex No. 2, No. 703-025	Do	Do.
Do	Proches No. 3, No. 705-025		Do.
Do	Proches No. 5, No. 706-025	Do	Do
2 Do	Proches No. 6. No. 707-025	Do. 4	Do.
Do	Prochex No. 7, No. 708-025	Do	Do. 14
Do	Prochex No. 8, No. 709-025	Do	Do.
Du	Prochex No. 9, No. 710-025	Do	Do.
Do	Prochex No. 10, No. 711-025	Do	Do.
Do	Ia) No. 712-025)		10.
Bio-Resgents &	Prochex No. 11, No. 713-025	Vial: 25 ml.	Mar. 9, 197
Disgnostics, Inc	Bunchas Mr. 19 Mr. 814 00E	Contraction of the second sec second second sec	The state
Do	Proches No. 12, No. (19-025 C	7	100. Do
Do	Proches No. 14, No. 716-025	Do	Do.
Do	Procher No. 15, No. 717-025	Do	Do.
Do	Prochex No. 15, (Alternate	Do	Do.
Bo	Prother No. 16, No. 219-025	1 Do	Da
Do	Proches No. 18 - No. 721-025	Do	Do.
Do	Proches No. 19, No. 722-025	Do	Do.
Do	Prochex No. 20, No. 723-025	Do	Do,
Do	Ia) No. 712-025) Prochex No. 11, No. 713-025 Prochex No. 12, No. 713-025 Prochex No. 13, No. 715-025 Prochex No. 14, No. 716-025 Prochex No. 15, No. 717-025 Prochex No. 15, Alternate Formula) No. 718-025 Prochex No. 16, No. 719-025 Prochex No. 18, No. 721-025 Prochex No. 18, No. 722-025 Prochex No. 20, No. 723-025 Prochex No. 20, No. 723-025 Toxicology control urine-dried No. 6716-25.	Bottle: 25 ml.	June 25, 19
Do	Toxicology control serum-dried No.	Bottle: 10 ml.	Do.
Do	trol-dried No. 6736-25.	Bottle: 20 mi.	
	Urine Control II No. 695-425 Brinkmann Drug-Skreen Drug		
Ing	standard-set T No. 8505000-1		
Do	Brinkmann Drug-Skreen Drug standard-set I, No. 3505000-1. Brinkmann Drug-Skreen Drug standard-set II, No. 3505010-8.		
Buchler Instruments	Buffer sait-type I, barbital-sodium,	11. er e.	
Do	Buchler instrument huffer R-2	Vial: 36.36 grams	. Sept. 15. 19

Manufacturer or supplier	Product name and supplier's catalog number	Form of product	Date of application
Burroughs Wellcome Co	Lanoxitest beta digoxin radioim- munoassay kit with tritiated dig- oxin No. KT07.	Bottle: 125 ml	Nov. 16, 1972
California Bionuclear Corp.	Amobarbital-2-C-14, catalog No. 72077.	Screw cap vial: 50 uCi, 0.1, 0.5, and 1 mCi.	Jan. 8, 1975
Do	D-Amphetamine (propyl-1-C-14) sulfate, catalog No. 72078.	Do	Do.
Do	DL-Amphetamine (propyl-1-C-14) sulfate, catalog No. 72079.	Do	Do.
Do	Cocaine (methoxy-C-14) catalog No. 72182.	Do	Do.
Do	Meperidine (N-methyl-C-14) hydro- chloride, catalog No. 72508.	Do	Do.
Do	Mescaline (aminomethylene-C-14) hydrochloride, catalog No. 72512.	Do	Do.
Do	Methadone (heptanone-2-C-14) hy- drochloride, catalog No. 72516.	Do	Do.
Do	Methamphetamine (propyl-1-C-14) sulfate, catalog No. 72517.	Do	Do.
Do	Methyl phenidate (carbonyi-C-14) hydrochloride, catalog No. 72550.	Do	Do. Do.
Do	Morphine (n-methyl-C-14) hydro- chloride, catalog No. 72560.	Do	Do.
Do	Pentobarbital-2-C-14, catalog No. 72618.		Do.
Do	Secobarbital-2-C-14, catalog No. 72675.	Ampoule: 50 uCi, 0.1, 0.5, and 1 mCi.	
Chemed Corp. (Dearborn Chemical Division)	Zinc reagent No. 2, No. 704	Pillow: 10 mg. each	June 23, 197
Collaborative Research, Inc.	Kit to include: LSD antiserum No. Z-20; I-125-LSD-Polymer No. Z- 11; LSD standard.	Bottle: 1 and 2 dram	Nov. 14, 197
Clarkson Laboratory and Supply, Inc.	Hematoxylin stain, Mayer's No. S- 1302.	Gallon	Dec. 12, 197
Collaborative Research, Inc.	Radioimmunoassay of Tetrahydro- cannabinol.	Kit containing: Δ^{0} -THC antiserum $^{3}H-\Delta^{0}$ -THC antigen- Dextran coated char- coal ¹⁴ C Δ^{0} -THC stand- ard	Jan. 5, 197
Do	"H 4"-THC Antigen	Normal rabbit serum Vial: 1 ml.	Do. Do.
Do Cordis Laboratories	¹⁴ C HA ⁹ -THC Standard Barbital-acetate buffer, powder 709-317.	Package: 20 envelopes —10.65 grams per en-	July 27, 197
Do	CEP V No. 709-308	velope. Plate: 80 mm. x 100	Aug. 9, 197
Do	CEP V No. 709-328	mm, x 2.2 mm. Plate: 40 mm. x 80 mm. x 2.5 mm.	Do.
Do Do	CEP VII No. 709-323 CEP V No. 709-338	Do Plate : 40 mm. x 80 mm.	Do. Do.
Do	CEP VI No. 709-309	x 2.5 mm Plate: 80 mm. x 100 mm. x 2.2 mm.	Do.
Cordis Laboratories	CEP VI No. 709-329	Plate: 40 mm. x 80 mm. x 2.5 mm.	Aug. 9, 197
Do Do	CEP VI No. 709-839 CEP plate-amebiasis testing 10 test	Do	Do.
Do	No. 730-271. CEP plate-amebiasis testing 40 test		Do.
Do	No. 730-274. Counterelectrophoresis, plates CEP	Package: 5 plates-18 ml. per plate.	Do.
Do	I 709-804. Counterelectrophoresis, plates CEP		Do.
Do		Do	Do.
Do	III 709-306. Counterelectrophoresis, plates CEP IV 709-307.	Do	Do.

Manufacturer or supplier	Product name and supplier's catalog number	Form of preduct	Date of spplication
Do	Counterelectrophoresis, plates CEP I 709-324.	CONTRACTORY DISTO	•** •* Do. *****
. Do	Counterplastrophoneric plater CER	Da	Do.
Do	II 709-325. Counterelectrophoresis, plates CEP III 709-326.	Do	Do.
Do	Counterelectrophoresis, plates CEP	Do	Do.
Do	IV 709-327. Counterelectrophoresis (CEP) Plates for Trichinosis Testing.	mm x 80 mm x 2.5	June 16, 1978
De	GVB#+ buffer: 758-087	Bottle: 50 ml.	Aug. 9, 1972
Do Do	Glucose-GVB ¹ / buffer, 753-036 EDTA (0.014AM)-GVB buffer,	Bottle: 50 ml.	Do. Do.
Do	GVB ³⁺ buffer: 753-087 Glucose-GVB ¹ / buffer; 753-036 EDTA (0.014AM)GVB buffer; 753-034. EDTA (0.01M)GVB buffer; 753-031.		Do
Da	5X Isotonic veronal buffer	Bottle : 1,000 ml	Do
Custia Nuclear Corp.	FERRONEX Kit No. 00250	Vial: 8 ml.	
Disconting The	Diaman Buffen No 65	Bottle: 1 gal.	May 7, 1975
Do	DiAgAu Buffer, No. 65 DiAgAu Plates, No. 50 DiAgAu Plates, No. 55	Plate: 16 ml.	Do.
		Colorenter Witch and and	June 12, 197
Do	T_4 Antigenum	Do	100. ¹
Do	125 I T-3 125 I T-3 125 I T-4 125 I T-4 Goat Anti-Rabbit Gamma Globulin	Do	De
Do manufacture	Goat Anti-Rabbit Gamma Globulin	Do	UDG.C.
Dow Chemical Co.	Iodine-125 Trilocolhyronine	VIAI: ZV.D MIL.	May 22, 197
D -	A - Alémile John and a Contine 1	Do	
Do Do	ANSA Builter Lyophilized	Do	Do
D9	Activated charcoal, T3 RIA	Do Vial: 3 g	May1, 197
Paul B. Elder Co.	Anturnootayronne Serum Lyophilized ANSA Buffer Lyophilized Dextran Lyophilized Activated charcoal, T3 RIA Fisher body heat indicator 338° F Tempilag	Bottle: pint Glass bottle: 16 fL oz	July 80, 197
Electro-Nucleonics Labora-	"338" F' Tempilaq Morphine (3R), List No. 4005 ' "Morphine Positive Control, List	Glass vial: 5 ml.	June 20, 197
Do	Morphine Positive Control, List No. 4006. Dextran Coated Charcoal Solution	Glass vial	Do.
Environmental Chemical Specialties, Inc.	Dextran Coated Charcoal Solution TBG-Radiothyroxine Solution	Bottle: 1,000 ml	
Do	TBG-Radiothyroxine Solution	Do	
Fisher Scientifio Co.	Electrophoretic buffer No. 1, pH 8,60, ionic strength 0.05, catalog No.E-1. Electrophoretic buffer No. 2, pH	Facket: 12.14 grams	. Oct. 27, 197
- ಸೌರ್ವೇಶ ಭೇಗಿಸುವುದು ಭೇಗಿಸುವುದು ಭೇಗಿಸುವುದು ಭೇಗಿಸುವುದು ಭೇಗಿಸುವುದು ಭೇಗಿಸುವುದು ಭೇಗಿಸುವುದು ಭೇಗಿಸುವುದು ಭೇಗಿಸುವುದು ಭೇಗಿ	No.B-1		. UN D e
Do	8.60, ionic strength 0.075, catalog	1	. Do.
Time Fahrendenfan	DCT No. 9 080	. Bottle: 125 ml	Apr 16. 197
PIOW LEDOFACOTICE	CEP Buffer No. 3-083	Bottle: 125 ml.	Do.
Do, C.	CEP Plate No. 5-076	Plate: 20 ml.	Do.
Do	DGV No. 3-080 CEP Buffer No. 3-083 CEP Plate No. 6-076 Merthiolate No. 6-088B Barblione acetate buffer for else	Bottle: 20 ml.	Do. June 21, 197
SLOID - FRY	Merthiolate No. 6-088B Barbitone acetate buffer for elec- trophoresis code No. BR 11g. Complement fixation test, diluent tablets code No. BR 16. Drug Standard Set, No. 51910	Bottle = 100 tablets	. Do.
Gelman Instruments Co.	brug Standard Set, No. 51910	Set: 3 vials of 2 ml.	Apr. 6, 191
Do	Drug Control Set, No. 51911	Set: 3 vials of 50 ml.	Do.
Do	High Resolution, buffer-Tris Barbl tal buffer, No. 51104.	- Viai: 10 dr	Dec. 22, 19
Concel Diamontics	Int Duller, NO. 01194.	Vial 10 6 and 1 9 and	Autor 96 10
General Anaguestics	fas T. No. 86908		
Grand Island Biological Co.	Dextrose-Gelatin-Veronal Buffer Solution NDC No. 815-0566-1 and No. 815-0566-2.	Bottle: 100 ml. and 500 ml.	July 5, 197

Manufacturer or supplier	Product name and supplier's catalog number	Form of product	Date of application
Do	Complement Fixation Buffer Solu- tion; pH 7.3-7.4, NDC 011815 0247 I.	Bottle:'1 Liter	Jan. 28, 197
Do Do	Diseragen, NDC 071815 1548 2	Vial: 100 ml	Nov. 21, 197 Do. Do.
	vine Albumin. Do Electrophoresis Buffer Solution, pH 8.6, NDC 011815 0245 1. I.E.P. Buffer Solution, pH 8.2,	Bottle: 1 Liter	Do. Jan. 28, ² 197 Do.
Do	NDC 011815 0246 1. Gibform Indicator Cells, NDC 011815 0220.	Viai: 40 ml	
Do	Gibform Adsorption Cells, NDC 011815 0225.	Vial: 20 ml.	Do.
Do Gugol Science Corp.		Glass bottle : 500 ml Visls : 20 ml., 90 ml., and 450 ml.	July 23, 197 Mar. 23, 197
lach Chemical Co.	pH 8.5 buffer powder pillows, No. 920-85.		Nov. 30, 197
Do	pH 8.3 buffer powder pillows, No. 898-98	Pillow: 1 gram each	Do.
Do	Zincover II powder pillows, No. 2917. Buffered substrate, glycerophos-	Vial: 0.855 gram per	Do.
	phate. Roe & Whitmore, pH 9.6, No. 20060.	104 101.	e servir
	Buffered substrate, glycerophos- phate, Shinowara, Jones & Reinhart, pH 10.9, No. 20063.	Vial: :0.925 gram per 100 ml.	Do.
Do	Reinnart, pri 10.9, No. 2005. Buffered substrate, glycerophos- phate, Shinowara, Jones & Reinhart-Stock, No. 20061.	Vial: 1.85 gram per 100 ml.	Do.
Do	Buffered substrate, glycerophos- phate, Shinowara, Jones & Reinhart, pH 5.0, No. 20062.	Vial: 0.925 gram per 100 ml.	Do.
Helena Laba.	Electra HR Buffer Catalog No.	packets per box.	"Dec. 28, 19
Do:	Electra B: Buffer Catalog No. 5017	Packet: 12.14 g, 10 packets per box. Packet: 18.21 g, 10	. Do.
Do Do	Titan III Agar Catalog No. 5023 . Titan IV IE Plate (amail)	packets per box. Vial: 2 ml. Package: plates, 1 by 3	Do. Do.
Do	Titan IV IE Plate (large)	in. Package: plates, 3 by 4 in.	Do.
Do 2	Titan IV IE Plate Kit	Kit: 12 small (1 by 3 in.) IE Plates, 1 box B ₁ Buffer.	Do.
	Titan IV IE Plate Kit	Kit: 10 large (3 by 4 in.) IE Plates, 1 box B ₁ Buffer.	Do.
Hoffman-La Roche, Inc.	Abusercen radio-immunoassay for morphine (1251), No. 43021.		Sept. 27 19
Do	Abuscreen radio-immunoassay for morphine (³ H), No. 43016. Abuscreen Radioimmunoassay for	Vial: 60 ml.	Do,
	Barbiturates ("H). Abuscreen Radioimmunoassay for	Vial: 60 ml. and 5 ml.	Ďo.
. Do	Barbiturates (191). Abuscreen Radioimmungassay for	Vial: 30 ml. and 500	Do.
Doູມ.	Barbiturates (125). Abuscreenim Mor-Barb Radioim- munoassay for Morphine-Barbi- turates	ml. Vial: 5 ml., 60 ml., and 100 ml. Bottle: 500 ml.	Dec. 27, 19

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Manufacturer or supplier	Product name and supplier's catalog number	Form of product	Date of application
Do	AbuscreentM Radio-immunoassay for Amphetamine.	Kit containing Vials of 5 ml., 30 ml., and 100 ml. and Bottle: 500 ml.	Jan. 14, 1974
Do	Abuscreen Radioimmunoassay for Morphine (1281).	Vial: 100 ml. and 500 ml.	Sept. 27, 1972
Do	Abuscreen Radioimmunoassay for Morphine (³ H).	Do	Do.
Do	Latex tube test kit for morphine:	Kit: 30 to 200 tests	Dec. 6, 1974
Do	TM Abuscreen radioimmunoassay for Methadone.	Kit containing vials of: 5 ml., 30 ml., and 100 ml., and bottle: 500 ml.	June 17, 1974
Do	TM Abuscreen radioimmunoassay for Methaqualone.	Do	Do.
Hyland Division Travenol Laboratories, Inc.	Agar gel plates No. 3008	Package: 8 plates-25 ml. per plate.	Aug. 31, 1971
Do	Agar gel plates No. 3016	Package: 10 plates-25 ml. per plate.	Do.
Do	Agar gel plates No. 8068	Do .	Do.
Do	Buffer No. 8017	Vial: 250 ml.	Do. Do.
Do	Buffer No. 8069 Diluting fluid No. 8400	Vial: 10 ml.	Do.
Do	Partial thromboplastin liquid No. 8481.	Vial: 0.1 ml.	Do.
Do	PTC reagent dried, No. 8497	Vial: 1 ml.	Do.
Hyland Division Travenol Laboratories, Inc.	Supplemental urine clinical chem- istry control, dried, No. 0402	Vial : 25 ml.	Aug. 81, 1971
Do	and No. 0521. Partial thromboplastin, dried, No. 8491.	Vial: 1 ml. and 5 ml	Do.
Do	Agar gel plates, No. 8794	Plate: 25 ml.	Aug. 1, 1972
Do	Buffer, No. 8793	Vial: 250 ml.	Do.
Do	Toxicology serum control, dried, No. 0541.	Vial: 10 ml	Oct. 26, 1972
Do	Toxicology urine control, dried, No. (542.	Do	Do.
Do	T-1	Vial: 20 ml	Jan. 13, 1976
Do Do	T-2 T-4		Do. Do.
Do	T-5	Do	Do.
Do	T-6	Vial: 20 ml.	Do.
Do	T-7	Do	Do.
Do Do	T-8 T-9	Vial: 50 ml.	Do. Do.
Do	T-10	Do	Do.
Do	T-11	Do Vial: 20 ml.	Do.
Do	T-12		Do.
Do	T-14 T-15	Vial: 50 ml.	Do. Do.
Do	T-16	Vial: 20 ml.	Do.
Do	T-18	Vial: 50 ml.	Do.
Do	T-20	Do Vial: 5 ml.	Do.
Do Do	TC-1	VIAI : 5 ml.	Do. Do.
ICL Scientific	EIQ Intensifier Diluent 1	Bottle: 7.6 gm.	Feb. 26, 1975 Do.
industrial Biological Laboratories, Inc.	DGV solution	Vial: 100 cc	Dec. 28, 1971
instrumentation Labora- tory, Inc.	Tris-Barbital Buffer No. \$3205	Vial: 12 dram	Feb. 21, 1971
Do	Barbital Buffer (B-2) No. 33206	Do	Do.
Do Do	EDTA-Barbital Buffer No. 33207 Barbital-Acetate Buffer No. 33208	DoDo	Do. Do.
nolex Corp. Biomedical Division.	Barbitone Acetate Buffer, Product Code 71-161-01.	Bottle: 125 gm	May 29, 1974
Do	Barbitone Acetate Buffer with Cal- cium Lactate, Product Code 71-162-01.	Do	Do.

Manui	facturer or supplier	Product name and supplier's catalog number	Form of product	Date of application
Do		Barbitone C.F.T. Diluent, Product Code 63-163-10.	Bottle: 100 tablets	Do.
	Delavau., Inc., and heta Corp.	Allobarbital No. FP305	Vial: 2 ml.	Apr. 10, 197
		Amobarbital No. FP313	Do	Do.
Do		Amobarbital No. FP313 Amphetamine No. FP604	Do	Do.
		Anileridine No. FP203	Do	Do.
Do		Aprobarbital No. FP306	Do	Do.
		Barbital No. FP314	Do	Do.
		Butabarbital No. FP815	Do	Do.
Do		Butalbital No. FP807	Do	Do.
		Chloral Betaine No. FP502	Do	Do.
	***	Chloral Hydrate No. FP501		Do.
		Cocaine No. FP601	Do	Do.
		Codeine No. FP102	Do	Do.
		Cyclobarbital No. FP308 Diphenoxylate No. FP205	Do	Do.
Do		Diphenoxylate No. FP205	Do	Do.
		Dyhydrocodeine No. FP108	Do	Do.
		Ethchlorvynol No. FP508 Ethylmorphine No. FP106	Do	· Do.
				Do.
		Fentanyl No. FP211	Do	Do.
Do		Glutethimide No. FP404	Do	Do.
<i>D</i> 0		Heptabarbital No. FP309 Hexobarbital No. FP303	Do	Do.
		Hydrocodone No. FP107	Do	Do.
		Hydromorphine No. FP103	Do	Do. Do.
		Levorphanol No. FP208	Do	Do. Do.
Do	**************	Marker Mixture No. FPM-104	Do	Do.
Do		Marker Mixture No. FPM-201	Do	Do.
		Meperidine No. FP201	Do	Do.
		Mephobarbital No. FP301	Do	Do.
		Meprobamate No. FP402	Do	Do.
Do	·	Methadone No. FP206	Do	Do.
Do		Methamphetamine No. FP603	Do	Do.
Do		Metharbital No. FP302	Do	Do.
Do	***************	Methohexital No. FP304	Do	Do.
Do		Methylphenidate No. FP605	Do	Do.
Do		Monthly Urine Test No. FPM-108	Do	Do.
	Delavau Co., Inc., he Theta Corp.	Morphine No. FP101	Vial: 2 ml	Apr. 10, 1973
		Oxycodone No. FP109	Do	Do.
		Oxymorphone No. FP104	Do	Do.
Do		Paraldehyde No. FP506	Do	Do.
no 10		Pentobarbital No. FP318 Phenazocine No. FP213	Do	Do.
100		Phenametrazine No. FP213	Do	Do.
		Phenobarbital No. FP320	Do	Do.
		Piminodine No. FP202	Do	Do.
	*******	Probarbital No. FP319	Do	Do.
			Do	Do. Do.
		Talbutal No. FP311	Do	Do. Do.
D		Thiamylal No. FP322	Do	Do.
		Thiopental No. FP321	Do	Do. Do.
		Vinbarbital No. FP312		Do.
		Weekly urine test (FDA) No. FPM- 101.	Do	Do.
Do	·····	Weekly urine test (States) No. FPM-102.	Do	Do.
	, 	Test mixture TM No. 1	Do	June 19, 197
De	*****	Test mixture TM No. 2	Do	Do.
Do		Test mixture SM No. 1	Do	Do. Do.
Do		spectively was alve a meansamene		De.
Do Do		Test mixture SP No. 1		
Do Do Do	****	Test mixture SP No. 1 Test mixture SM No. 2	Do	Do.
Do Do Do Do		Test mixture SM No. 2	Do	Do. Do.
Do Do Do Do		Test mixture SM No. 2	Do	Do,
Do Do Do Do Do		Test mixture SM No. 2 Test mixture SP No. 2 Test mixture SM No. 3 Test mixture SP No. 3	Do Do Do Do Do	Do. Do.
Do Do Do Do Do Do		Test mixture SM No. 2		Do.

Manufacturer or supplier	Product name and supplier's catalog number	-		
Kailestad Labs, Inc.	Osmotect Buffer No. M 101		May 17, 197:	
Do	Buffer No. C135 Osmotect Agar Gel Plate Kit No. M 100.	Wint. 7 Jun	Do. Do.	
oderle Laboratories Divi- sion of American Cyan-	DGV buffer, 5x No. 2606-37	Vial : 20 ml.	Nov. 19, 1971	
amid Có. Do	Serum toxicology control drugs A, No. 2040-69.	Vial: 10 ml.	Do.	
Do	Abnormal urine control, No. 2921- 80.	Vial: 25 ml.	Do.	
Do	Urine toxicology control drugs I,			
Do	Droficiency, No. 2551-61.	·.	Mar. 13, 1972	
Do	Urine toxicology control drugs 2- barbiturates, No. 2952-61.	Do	Do.	
De	Urine toxicology control drugs 2	Du	Do.	
Do	2953-61. Urine toxicology control drugs 3		Do.	
Do	Urine toxicology control drugs 8- amphetamines, proficiency No. 2955-61.	Do	Do.	
Do	Urine toxicology control, drugs 4-	Do	De.	
Do	Urine toxicology control, druga 4	Do	Do.	
Lederle Laboratories	61. Urine drug check kit No. 2953-93 to include: UDC 1 No. 2959-33. UDC 1a No. 2959-33. UDC 2 No. 2960-38 UDC 3 No. 2961-38 UDC 4 No. 2962-38 UDC 5 No. 2963-38 UDC 6 No. 2965-38 UDC 7 No. 2965-38 UDC 8 No. 2965-38 UDC 9 No. 2965-38 UDC 9 No. 2965-38 UDC 9 No. 2965-38 UDC 9 No. 2965-38 UDC 10 No. 2968-38 UDC 10 No. 2968-38	Bottle: 25 ml	Apr. 4, 197	
Do	to include: UDC 1 No. 2959-38. UDC 1a No. 2979-38	Do	. Do. :	
Do	UDC 2 No. 2960-38	Do	. Do.	
Do Do	UDC & No. 2961-38		- 10. Do	
Do	UDC 5 No. 2963-38		Do.	
Do	UDC 6 No. 2964-38		Do.	
Do	UDC 7 No. 2965-38	Do	Do.	
Do	UDC 8-No. 2966-38		Do.`	
Do Do Do Do Do Do Do Do Do	UDC 9 No. 2967-38		_ Do.	
Do	UDC 10 No. 2968-38	Do	. Do.	
Do	UDC 10 No. 2903-38 UDC 11 No. 2969-38 UDC 11 No. 2969-38 UDC 12 No. 2970-38 UDC 13 No. 2971-38 UDC 14 No. 2972-38 UDC 14 No. 2972-38 UDC 15 No. 2973-38	· ····· 20 -·····	. Do.	
1/0	. UDC 11 No. 2969-38		. Do.	
D0	UDU 12 NO. 2970-38	<i>10</i>	- Do.	
Do	. UDU 14 NO. 29/1-08	/0	Do	
10	LIDC 15 No. 9072 34		Do. Do.	
Do	UDC 15a No. 2981-38	Do	. Do.	
Do	UDC 16 No. 2974-38		Do.	
Lederle. Laboratories	_ UDC 17 No. 2975-38	_ Bottle: 25 ml	Apr. 4, 197 Do.	
Do Do	UDC 19 No. 2977-38	Do	Do.	
LKB Instruments, Inc	 Barbital Buffer pH 8.6, No. LKB 5104-180. 	- Vial: 290 ml.	Jan. 3, 197	
Mallard, Inc.	- High resolution buffer-tria ban bital buffer No. 51104.	•.		
Mallinckrodt Chemical Works.	Res-O-Mat ETR solution	Vial: 1½ dram Do	_ Feb. 17, 197	
Do	_ Res-O-Mat ETR Solution	Buttle: 16 oz. and in	- Aug. 28, 197	
Do	Res-O-Mat T4 Solution	Do	Do.	

Maanfacturer or supplier	Product name and supplier's catalog number	Form of product	Date of application
fallinckrodt	RIA-MAT Circulating T3 1125 Kit.	Kit containing the fold	Jan. 28, 1974
	Cat. No. 501:	lowing :	
5 ⁻²	RIA-MAT T3 Buffer	Bottle: 100 ml.	
	RIA-MAT T3 Antiserum	Vial: 2.5 ml.	
and the second	RIA-MAT T3 Reaction Vial RIA-MAT T3 Standard 0 ng/ml	Vial: 1 ml. Vial: 1.5 ml.	· · · ·
	RIA-MAT T3 Standard 0.5	Vial: 1.5 ml.	
144 (A. 1997)	ng/ml.		4
· · · · · · · · · · · · · · · · · · ·		Vial: 1.5 ml.	
and a second s	ng/ml.		
		Vial: 1.5 ml.	
	ng/ml.	· · · · · · · · · · · · · · · · · · ·	
and the second second	RIA-MAT T3 Standard, 6.9	Vial: 1.5 ml.	3.85
••••	RIA-MAT T4 I-125 kit	Kite containings 100	Ann 9 1071
و با با با با با با با با		tests and 250 tests.	
laterials & Technology	Carboxymethylmorphine Sensitized	Vial : 50 ml	May 3, 1973
Systems, Inc.	REC.		
Do,	Carboxymethyl-morphine bovine	Vial: 8 ml.	Do.
	albumin.		
Do	albumin. Ectonine Sensitized RBC 5-ethyl-5-(1-carboxy-n-propyl) bar- bituric acid Sensitized RBC.		Do.
Do	5-ethyl-5-(1-carboxy-n-propyl) bar-	Do	Do.
	bituria acid Sensitized RBC. S-ethyl-5-(1-carboxy-n-propyl) bar- bituric acid. S-ethyl-5-(1-carboxy-n-propyl) bar-	• • •	
Do	5-ethyl-5-(1-carboxy-n-propyl) bar-	Vial: 8 mi. and 10 mi	Do.
Do	5-ethyl-5-(1-carboxy-n-propyl) bar-	3/1-1-0-1	Do.
D O	bituric acid bovine serum al-	V INI C MMr annaan	100,
. 4 1	bumin or rabbit serum albumin.	2.3 :	
Da	Ecgonine bovine serum albumin or		Do.
	rabbit serum albumin.		20.
Do	Tropinecarboxylic acid	Vial: 8 ml. and 10 ml	Do.
Do	Morphine standard	Vial: 10 ml	July 17, 197
Do	Morphine-urine standard	Vial: 25 ml	Do.
Do Do	Ecgonine-urine standard Barbiturste-urine standard	Do	Do.
Do	Benzoyi Ecgonine	. Do Vial: 25 mg. and 100	Do. Apr. 18, 197
		mg.	and the set
	Benzoyl Ecgonine-Urine Standard_	Vial: 25 ml. 3 mcg./ml.	Do.
Do Do	Rentual Francine Linne Standard	Visl 75 mor	Do,
dadi ing ka	Lyophilized	14M - 17 - 17 - 17	
Do	Cocaine-Urine Standard Cocaine-Urine Standard Lyophi-	Vial: 25 ml. 3 mcg./ml.	Do.
Do	Cocaine-Urine Standard Lyophi-	Vial: 75 mcg.	Do.
	lized	A	
Do	Methadone-Urine Standard	Vial: 25 ml. 3 mcg./ml.	Do.
Do	"Methadone-Urine Standard Lyophi-	-Vial: 75 mcg.	Do.
Da	Disco. Phenobarbital-Urine Standard Ly- ophilized. art. Secobarbital-Urine Standard Secobarbital-Urine Standard Lyo- philized	Vial: 25 ml. 5 mcg./ml.	Do.
Do	Phenobarbital-Urine Standard Ly-	Vial: 125 mcg.	Do,
Do	Secolarbital-Urine Standard	Vial . 25 ml. 8 mcg./ml.	Do.
Do	Secobarbital-Urine Standard Lyo-	Vial: 125 mcg.	Do.
	Main addite		
Dolig Hilliganoon	Amobarbital-Urine Standard	Vial: 25 ml. 5 mcg./ml.	Do.
Do 13	Amobarbital-Urine Standard Lyo-	Vial: 125 mcg	Do.
	philized.	·· · · · · · · · · · · · · · · · · · ·	
MCI Biomedical	IEP buffer; pH 8.2, 0.04 ionic strength.	Package: 6.510 grams_	"Aug. 28, 197
ALAD Diagnostics	T-3 test fas T. No. L6902	Vial: 3/1" x 1 13/16"	May 31, 1972
Medi-Chem, Inc.	. Thymol-Barbital Buffer Concen-	Vial: 10 ml	July 11, 197
Level de Mar	Thymol-Barbital Buffer Concen- trate pH 7.55. Thymol-Barbital Buffer Concen- trate pH 7.8		- n- ¹
Do	Thymol-Barbital Buffer Concen-	Do	Do.
	. Secobarbital Standard 10 mg. per- cent, Cat. No. 250.		Feb. 22, 197

Manufacturer er supplier	Product name and supplier's catalog number	Form of product	Date of application
Meloy Laboratories	Counterclectrophoresis . Pistes, G- 301.	Plates: 10 determina- tions.	Sept. 5, 1973
	Immunoelectrophoresis Plates, G- 201.		Do.
Do	Immunostatim T3 Kit, No. K130 _	Cardboard box: 8%" x 5¼" x 2¼".	July 7, 1973
	Immunostatus T4 Test Kit, No. K140.	Cardboard box: 8%" x	Do.
Lederle Laboratories	UDC 17 No. 2975-38	Bottle: 25 ml.	Apr. 4, 1973 Do.
Do2	UDC 19 No. 2977-38 UDC 20 No. 2978-38	Do	Do. Do.
LKB Instruments, Inc.	UDC 17 No. 2975-38 UDC 17 No. 2976-38 UDC 18 No. 2976-38 UDC 19 No. 2977-38 UDC 20 No. 2978-38 Barbital Buffer pH 8.6, No. LKB- 5104-180.	Vial: 290 ml.	Jan. 3, 197
Mallard, Inc.	5104-180. High resolution buffer-tris bar- bital buffer No. 51104.	Vial: 1½ dram	Dec. 22, 197
Mallinckrodt [†] Chemical Works.	Res-O-Mat ETR solution		
Do	Res-O-Mat T4 solution		Do
Do.,	Res-O-Mat T4 solution Res-O-Mat ETR Solution	Bottle: 15 oz. and im- perial gallon.	Aug. 28, 197
Do	Res-O-Mat T4 Solution	Do	Do.
Mallinekrodt	RIA-MAT Circulating T3 1125 Kit, Cat. No. 501: RIA-MAT T3 Buffer RIA-MAT T3 Antiserum	Kit containing the fol- lowing:	Jan. 28, 197
	RIA-MAT T3 Buffer	lowing: Bottle: 100 ml. Vial: 2.5 ml. Vial: 1 ml.	
	RIA-MAT T3 Reaction Vial RIA-MAT T3 Standard 0 ng/ml	7 1004 8 . 4 16161	
	RIA-MAT T3 Standard 0.5	Vial : 1.5 ml. Vial : 1.5 ml.	
. [.] .	ng/ml. RIA-MAT T3 Standard 1.0 ng/ml.	Vial: 1.5 ml.	
		Visl: 1.5 ml.	
·· »»· ·	RIA-MAT T3 Standard 6.0	Vial: 1.5 ml.	
an a	RIA-MAT T4 I-125 kit	Kits containing: 100 tests and 250 tests.	Apr. 3, 197
Mataviala & Technology	Carboxymethylmorphine Sensitized	Vial: 50 ml	May 3, 197
Do	Carboxymethyl-morphine	Vial: 8 ml.	Do.
Do	RBC. Carboxymethyl-morphine Carboxymethyl-morphine bovine serum albumin or rabbit serum	Do	Do.
		Do	De
Do Do	alkumin. Ectonine Sensitized RBC 5-ethyl-5-(1-earboxy-n-propyl) bar- bituric acid Sensitized RBC.	Do	Do. Do.
Do	o-scual-o-(I-carboxà-u-brobàr) par-	Vial: 8 ml. and 10 ml.	
Do	biturie scid. 5-ethyl-5-(1-carboxy-n-propyl). bar biturie. scidbovineserumsl-	Vial : 8 ml	Do.
Do	bumin or rabbit serum albumin. Ecgonine bovine serum albumin	Do	Do.
Do	or rabbit serum albumin. Tropinecarboxylic acid	Vial: 8 ml. and 10 ml	Do.
Do	Morphine standard	Vial: 10 ml.	July 17, 197
Do Do	Morphine-urine standard	.Vial: 25 mlDo	Do. Do.
Do	Barbiturate-urine standard	Do	Do.
Do	Barbiturate-urine standard Benzoyl Ecgonine	Vial: 25 mg. and 100 mg.	
Do Do		Vial: 25 ml. 3 mcg./ml.	
i Do		Vial: 25 ml. 3 mcg./ml	. Do.
	Cocaine-Urine Standard Lyophi- lized.	Vial: 75 mcg.	Do.
	Methadone-Urine Standard Methadone-Urine Standard Lyo-	Vial: 25 ml. 3 mcg./ml	Do. Do.

Manufacturer or supplier	Product name and supplier's catalog number	Form of product	Date of application
Do Do	Phenobarbital-Urine Standard Phenobarbital-Urine Standard Ly- ophilized.	Vial: 25 ml. 5 mcg./ml. Vial: 125 mcg.	Do. Do.
Do	Secobarbital-Urine Standard Secobarbital-Urine Standard Ly- ophilized.	Vial: 25 ml. 5 mcg./ml. Vial: 125 mcg.	Do. Do,
Do	Amobarbital-Urine Standard Ly- ophilized.	Vial: 25 ml. 5 mcg./ml. Vial :125 mcg.	Do. Do.
MCI Biomedical	IEP buffer; pH 8.2, 0.04 ionic strength.	Package: 6.510 grams_	Aug. 28, 1972
MEAD Disgnostics	T-3 test fas T, No. L6902 T-4 test fas T ₁ , No. L6905	Vial: 1/2" x 1 13/16" Vial: 1/2" x 1 13/16"	May 31, 1972 Do.
Medi-Chem, Inc.	Thymol-Barbital Buffer Concen- trate pH 7.55.		July 11, 1974
Do	Thymol-Barbital Buffer Concen- trate pH 7.8.	Do	Do.
Medical Chemical Corp	Secobarbital Standard 10 mg. per- cent, Cat. No. 250.	Bottle: 120 cc	Feb. 22, 1974
Meloy Laboratories	Counterelectrophoresis Plates, G- 301.	Plates: 10 determina- tions.	Sept. 5, 1973
Do	Immunoelectrophoresis Plates, G- 201.	Plates: 6 per Unit	Do.
Do	Immunostatim T3 Kit, No. K130	Cardboard box: 85%" x 51%" x 21%". Cardboard box: 85%" x	
Do	Immunostatim T4 Test Kit, No. K140.	Cardboard box: 8% x 51/4" x 21/4".	D o,
Purex Laboratories, Inc	tract, 1,000 pnu/cc.	Vial: 2 cc	•
Do	Cannabis sativa, allergenic ex- tract, 20,000 pnu/cc.	Vial: 50 cc	Do,
Ortho Diagnostica	Activated Thromobo FAX No. 721000.	Bottle : 3.2 ml	Sept. 21, 197
Do	Hapindex, agar gcl plate, No. 74000.	Plate: 43 ml. per plate	Do.
Do	tion control.		
Do	Ortho HAA positive control No. 740100.	Vial: 1 mg	Mar. 27, 197
Ortho Diagnostics Oxford Laboratories Do	Ortho Control Urine II, No. 9040 StaT4 Adsorbent? Catalog No. 991. StaT4 Adsorbent, Catalog No. 992.	Vial: 25 ml. Lyophilized Bottle: 95 ml.	-
Oxy Metal Industries Corp. Regis Chemical Co Do	Compound N Solution	Steel drum: 55 gal Vial: 5 mi Vial:	Oct. 1, 197
	Group A Group B Group C Group D Group E	5 ml. Do Do Do 5 mi.	
Schering Corp Schwarz/Mann Division.	Hepaquik D L-amphetamine sulfate C14	Vial: 9 dram and plate. Flask: 0.05 mc, 0.1 mc,	
Becton Dickson and Co. Do	sterile aqueous solution. D-amphetamine sulfate C14 sterile aqueous solution.	0.5 mc, 1.0 mc.	Do.
Do	L-amphetamine sulfate C14 sterile aqueous solution.	Do	Do.
Do Do Do	Secobarbital 5 C14 Secobarbital 2 C14 Barbital buffer salt mixture, No. 0752-04 and No. 0752-07.	Do Do Vial: 50 cc.	Do. Do. Nov. 4, 197
SGA Scientific Corp Do	Barbital-acid buffer salt, No. 1173.	Bottle : 4 oz.	Do. Do.

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lanufacturer or supplier	Product name and supplier's catalog number	Form of product	Date of application
Do	Barringer & Woodard buffered	Vial: 0.73 gram per 15	Sept. 15, 1971
Do	Buchler instrument buffer B-2 double strength, pH 8.6, 0.075 m No. 93834.	x 45 mm. vial. Vial: 36.36 grams	Do.
	double strength, pH 8.6, 0.075	THE OVICE STORES	N U.
2	m No. 93834.		
Do	Buffer barbital, pH 8.8, No. 7691.	Vial: 11.76 grams per	Do.
	_ Buffer salt-barbital acetate, mix-	10 dram vial. Vial: 14.7 grams per	Do.
	ture pH 8.6, No. 3787.	23.5 Y X0 mm, VILL	
• Do	_ Buffer salt mixture pH 8.3, No. 7644.	Vial: 17.85 grams per 29.5 x 80 mm. vial.	Do.
Do			Do.
	Buffer salt mixture Spinco B-1, pH 8.6, 9.05 ionic strength, No.	Vial: 12.12 grams per 29.5 x 80 mm. vial.	
	3947. Buffer sait mixture Spinco B-2,	32-1. 10-10	
Do	pH 8.6, 0.075 ionic strength, No.	Vial: 18.18 grams per 29.5 x 80 mm. vial.	Do.
· · · ·	8948.	·· · · ·	. 19"
_ Do	Buffered barbital, sodium chloride,	Vial: 14.7 grams per vial.	Do.
Do	pH 7.5, No. 646-7. Buffered substrate glycerophos-	Viai: 0.924 gram per	Do.
Do	phate Bodansky No. 23681.	15 x mm. vial.	
	. SUITERED VERODAL DIS 7.3. NO.	Vial: 16.48 grams per vial.	Do.
Do	Gilerees & Davis buffered sub- strate, No. 23701.	Vial: 1.228 grams per	Do.
Reg I is a for a figure	Strate, No. 23701.	15 x mm. vial.	
Do annananananan	King & Armstrong buffered sub- strate; No. 23721. Ros & Whitmore buffered sub-	Vial: 1.14 grams per 15 x 45 mm. vial.	Do.
Do	Roe & Whitmore buffered sub-	Vial: 0.854 gram per	Do.
Do	strate, No. 23686. 	Vial: 18.18 grams per	Do.
	ture B-2. No. 93953	10 dram vial.	
Do	Iered substrate, No. 23733.	Vial: 0.945 gram per 	Do.
Do	. Thymol barbital buffer McLagan	Vial: 1.256 grams per	Do.
Do	Thymol buffer 100 ml-100 mg.,	15 x 45 mm. vial. Vial: 0.964 gram per 15 x 45 mm. vial.	Do.
Do the first	Huerga & Pepper, No. 29959. Thymol buffer pH 7.8, MacLagan,	15 x 45 mm. vial.	
	Ng. 29949.	VIAI.	Do.
Do	. Thymol buffer pH 7.55 Mateer,	Vial: 0.96 gram per 15	Do. ``
Do	No. 29951. _ Thymol turbidity test set. No.	x 45 mm. vial. Packet: 1 gram	Nov. 4. 1971
	3105.		
Do		Vial: 0.514 gram per vial.	Sept. 15, 1971
Do	_ Adenosine phosphate substrate No.	Botule: 4 oz	July 25, 1973
Do	675-1. Glycerophosphate substrate No.	The.	Da
1/0	675-2.	· · · · · · · · · · · · · · · · · · ·	Do.
Do	_ Glycerophosphate substrate No.	Do	Do.
GA Scientific Corp		Bottle: 25 ml and	Aug 6 107
	280-2.	100 ml.	AUG. 4, 121
. Do	_ Mayer's hematoxylin solution No.	Do	Do.
Do	MHS-1.	Wale a mi	
No	SGOT Single Assay Visi, No. 55-1.	Viki: 3 mi	May 29, 1973
Do	SGOT Assay Vial. No. 55-5	Vial: 15 ml	Do.
Do	SGOT 10 Assay Vial, No. 55-10	Vial: 30 ml.	Do. Do.
	IP.		
Do	SGPT 5 Assay Vial, No. 55-5P SGPT 10, Assay Vial, No. 55-	Vial : 15 ml.	Do.
. No			Do.
Do	SCOT Reagent No. 155-10 SGOT Reagent No. 155-100 SGPT Reagent No. 155-10P SGPT Reagent No. 155-10P	.Do	Do.
		Wint, \$60	· · · · ·
Do	SGUT Resgent No. 155-100	viai: 100 mi	Do. Do.

tries Re-agent kit catalog No. 8889- 007608. Tet Kit containing: 25 plastic tubes coated with anihody. Oct. 15, 197 T-3 Inc. Tet Kit containing: T-3 Inc. Oct. 15, 197 25 plastic tubes coated with anihody. Oct. 15, 197 F. R. Squibb & Sons Barbital Buffer Mixture for use with Gastrin Immutope Kit No. 095610. I vial of barbital buffer. Nov. 21, 197 Do AuSure II Barbital Buffer Pow- der, No. BT8209. Vial: 5.01 gm. July 28, 197 Do Barbital buffer mixture for use with discrin immutope Kit No. 093500. Vial: 6.055 gm. De 1, 197 Do Barbital buffer mixture for use with discrin immutope Kit No. 09350. Vial: 6 cc. July 30, 197 Do Thyrostat-4 Kit, Catalog No. 09125. To include: (a) Thyrostat-4 Standard Solu- vial (b) Tarprotat-4 Buffer Solution. Do Bottle: 60 ml. June 5, 197 Do Coninc. 64-9183 1,000 %g/glass ampul June 5, 197 Do. 200 Juse 64, 197 Do Coninc. 64-9183 1,000 %g/glass ampul Do. 200 Do. 21,000 wg/glass ampul Do. 22, 1971 Do Coninc. 64-9183 Do Do Do. 20, 21, 21, 21, 21, 21, 22, 21, 21, 21, 21	Manufacturer or supplier	Product name and supplier's catalog number	Form of product	Date of application
Tries Re-agent kit catalog No. 8889- 007668. Smith Kline Instruments, T-3 Inc. Uptake Disgnostic Test Test Kit containing: 25 plastic tubes coated with antibody. Oct. 15, 197 Smith Kline Instruments, T-3 Inc. Uptake Disgnostic Test Test Kit containing: 25 plastic tubes coated with antibody. Oct. 15, 197 E. R. Squibb & Sons			Vial: 30 ml. Vial: 100 ml.	
Smith, Kline Instruments, Uptake Diagnostic Test Test Kit containing: 25 plastic tubes costed with antibody. Standards. Oct. 15, 197. T-3 Inc. Barbital Buffer Mixture for use 09510.0 Standards. Nov. 21, 197. E. R. Squibb & Sons Barbital Buffer Mixture for use 09510.0 Nov. 21, 197. Nov. 21, 197. Do AuSure II Barbital Buffer Pow- der, No. 27920.0 Vial: 6 cc. Nov. 21, 197. Do Barbital buffer mixture for use 09510.0 Part Will Science Nov. 21, 197. Do Barbital buffer mixture for use 09550.0 Part Will Science Nov. 21, 197. Do Barbital buffer mixture for use with digoxin immutope kit No. 09350.0 Nov. Therestat-4 Kit, Catalog No. 09350.0 Do Feb. 26, 197. Do Thyrostat-4 Standard Solu- 09125. To include: (a) Thyrostat-4 Standard Solu- 09560.0 Vial: 7 ml. July 30, 197. Do Barbital Duffer Mixture for use with Digoxin Immutope Kit No. 09560.1 1,000 ug/glass ampul Userglic Acid Distribuinde Tar- trate, 04-9153 1,000 ug/glass ampul Do June 5, 137. Do Peliocin, 04-9161 Do Do Do Do Do Do Do Do Do		Re-agent kit catalog No. 8889-	Kit	April 17, 1975
1 vial of barbital buffer. Barbital Buffer Mixture for use with Gastrin Immutope Kit No. Vial: 5 cc. Nov. 21, 197. bo AuStre II Barbital Buffer Powder, No. 878209. Piate: 30 microliters Sept. 16, 197. bo AuStre II CEP Plate No. 878209. Piate: 30 microliters Sept. 16, 197. bo Barbital buffer mixture No. 09501. Parts and buffer mixture No. 09501. Vial: 6 cc. July 30, 197. bo Barbital buffer mixture No. 09501. Parts and buffer mixture No. 09503. Dec. 21, 197. bo Barbital buffer mixture for use with digoxin immutope kit No. 09125. To inclute: Yul: 5 cc. July 30, 197. bo Barbital Buffer Mixture for use with ligoxin immutope kit No. 09125. To inclute: Yul: 2.4 g. per 6 ml. Aug. 6, 197. bo Barbital Buffer Mixture for use with ligoxin immutope kit No. 09360. Vial: 2.4 g. per 6 ml. Aug. 6, 197. Do bo Bo Cocatine, 04-9183 1,000 ug/glass ampul _ Do. Do Do Do bo Pailocybin, 04-9190 How Pailocybin, 04-9172 Do Do Do Do Do			25 plastic tubes coated with antibody. Standards. 1 vial of radioactive	Oct. 15, 1975
with Gastrin Immutope Kit No. 09510. July 28, 197: der, No. B72209. July 28, 197: der, No. B72209. July 28, 197: der, No. B72209. Do July 28, 197: der, No. B72209. Do Do Barbital buffer mixture No. 09501. Do Do Do Barbital buffer mixture No. 09501. Uial: 6.065 gm Dee. 21, 197: July 28, 197: Do Thyrostat-4 Kit, Catalog No. 09125. To include:: (a) Thyrostat-4 Standard Solu- tion. Uial: 2.4 g. per 6 ml. Aug. 6, 187: with Digoxin Immutope Kit No. 09360. Supelco, Inc. Cocceine, 04-9188 Lysergic Acid Diebylamide Tar- 00 Do Do Do Do Do Do Do Do Do Do Do Do Do Do <td></td> <td></td> <td>1 vial of barbital buffer.</td> <td></td>			1 vial of barbital buffer.	
der, No. B72209. Plate: 30 microliters Sept. 16, 197. Do Barbital buffer mixture No. 09501. Vial: 6.065 gm. Dec. 21, 197. Do Barbital buffer mixture for use with digoxin immutops kit No. Vial: 6.065 gm. Dec. 21, 197. Do Thyrostat-4 Kit, Catalog No. Feb. 26, 197. July 20, 107. Do Thyrostat-4 Standard Solu- Vial: 7 ml. Feb. 26, 197. Do Barbital Buffer Mixture for use Vial: 2.4 g. per 6 ml. Aug. 6, 197. Yial. 60. Do Barbital Buffer Mixture for use Vial: 2.4 g. per 6 ml. Aug. 6, 197. Yial. 60. Do Coccaine, 04-9188 Do Do Jono %g/glass ampul Jone 5, 197. Do Lysergic Acid Diethylsmide Tar- Tarde, 04-9195. Do	E. R. Squibb & Sons	with Gastrin Immutope Kit No.	Vial: 5 cc	Nov. 21, 1972
Do Austre 11 CEP Plate No. 578209. Plate : 30 microiters Sept. 16, 197. Do Barbital buffer mixture No. 09501. Vial : 6.065 gm. Dec. 21, 197. Do Barbital buffer mixture for use with digoxin immutope kit No. 09350. Vial : 6.065 gm. Dec. 21, 197. Do Thyrostat-4 Kit, Catalog No. 0125. To include: Vial : 6 0ml. Dec. 21, 197. Do Thyrostat-4 Kit, Catalog No. 0125. To include: No. 0125. To include: No. 0125. Do Thyrostat-4 Buffer Solution. 015 Nyrostat-4 Buffer Solution. 019360. Bottle: 60 ml. Dottle: 60 ml. Dottle: 60 ml. Dottle: 60 ml. Dot. Dottle: 60 ml. Dot. Dot. Dottle: 60 ml.	E. R. Squibb & Sons, Inc.	der, No. B79209.		_
with algorin immutope kit No. 09350. Do Thyrostat-4 Kit, Catalog No. Feb. 26, 197. 09125. To include: (a) Thyrostat-4 Standard Solution. Vial: 7 ml. Feb. 26, 197. Do	Do	_ AuSure II CEP Plate No. B78209.	Plate: 30 microliters per well.	Sept. 16, 1971
09125. To include: To include: Bottle: Co ml. Source:	Do Do	 Barbital buffer mixture for use with digoxin immutope kit No. 	Vial: 6.055 gm Vial: 5 cc	Dec. 21, 1972 July 80, 1973
(b) Thyrostat-4 Buffer Solution. Bolle: 60 ml.	Do	09125. To include: (a) Thyrostat-4 Standard Solu-	•	Feb. 26, 1973
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Do	 (b) Thyrostat-4 Buffer Solution. Barbital Buffer Mixture for use with Digoxin Immutope Kit No. 	Vial: 2.4 g. per 5 ml.	Aug. 6, 1974
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Supelco, Inc.	- Cocaine, 04-9188	1,000 ug/glass ampul _	June 5, 1975
Do Amobarbital No. 04-9170 Ampule: 1 ml. Dec. 22, 197 Do Amphetamine No. 04-9165 Do Do Do Do Do Barbital No. 04-9171 Do Do Do Do Do Barbital No. 04-9171 Do Do Do Do Do Barbital No. 04-9175 Do Do Do Do Do Codeine No. 04-9175 Do Do Do Do Do Cyclobarbital No. 04-9176 Do Do Do Do Do Glutethimide No. 04-9176 Do Do Do Do Do Do Methadone No. 04-9163 Do Do Do Do Do Do Methamphetamine No. 04-9168 Do Do Do Do Do Do Methadone No. 04-9178 Do Do Do Do Do Do Methadone No. 04-9179 Do Do Do Do Do Do Do <td>Do</td> <td>trate, 04-9195.</td> <td>500 ug/glass ampul</td> <td></td>	Do	trate, 04-9195.	500 ug/glass ampul	
Do Amobarbital No. 04-9170 Ample: 1 ml. Dec.		Psilocin, 04-9190		
Do Amphetamine No. 04-9165 Do Do Do Do Aprobarbital No. 04-9171 Do Do Do. Do Barbital No. 04-9171 Do Do Do. Do Butethal No. 04-9172 Do Do. Do. Do Codeine No. 04-9172 Do Do. Do. Do Codeine No. 04-9173 Do Do. Do. Do Codeine No. 04-9175 Do Do. Do. Do Giutethimide No. 04-9176 Do Do. Do. Do Heptabarbital No. 04-9176 Do Do. Do. Do Methadone No. 04-9163 Do Do. Do. Do Methamphetamine No. 04-9178 Do Do. Do. Do Methamphetamine No. 04-9178 Do Do. Do. Do Methamphetamine No. 04-9179 Do Do. Do. Do Methamphetamine No. 04-9179 Do Do. Do. Do Methamphetamine No.	Do	Amobarbital No. 104-9170	Ampule: 1 ml.	Dec. 22, 1972
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Do '	Amphetamine No. 04-9165	Do	Do.
Do Butetehal No. 04-9172 Do Do Do Do Codeine No. 04-9161 Do Do Do. Do. Do Cyclobarbital No. 04-9175 Do Do. Do. Do Glutethimide No. 04-9175 Do Do. Do. Do Heptabrital No. 04-9175 Do Do. Do. Do Heptabrital No. 04-9176 Do Do. Do. Do Heptabrital No. 04-9162 Do Do. Do. Do Methadone No. 04-9163 Do Do. Do. Do Methamphetamine No. 04-9163 Do Do. Do. Do Methamphetamine No. 04-9178 Do Do. Do. Do Methamphetamine No. 04-9179 Do Do.	Do	- Aprobarbital No. 04-9171	Do	
Do Codeine No. 04-9161 Do Do Do Do Cyclobarbital No. 04-9175 Do Do. Do. Do Glutethimide No. 04-9175 Do Do. Do. Do Heptabarbital No. 04-9176 Do Do. Do. Do Heroin No. 04-9162 Do Do. Do. Do Methadone No. 04-9177 Do Do. Do. Do Methadone No. 04-9163 Do Do. Do. Do Methadone No. 04-9168 Do Do. Do. Do Methadone No. 04-9168 Do Do. Do. Do Methadone No. 04-9168 Do Do. Do. Do Methamphetamine No. 04-9178 Do Do. Do. Do Morphine No. 04-9179 Do Do. Do. Do. Do Pentobarbital No. 04-9179 Do Do. Do. <td< td=""><td>Do</td><td>Butetehal No. 04-9172</td><td>D0</td><td></td></td<>	Do	Butetehal No. 04-9172	D0	
Do Cyclobarbital No. 04-9175 Do Do Do Glutethinide No. 04-9178 Do Do Do Do Heptabarbital No. 04-9178 Do Do Do Do Heroin No. 04-9162 Do Do Do Do Do Heroin No. 04-9163 Do Do Do Do Do Methadone No. 04-9163 Do	Do	Codaina No. 01-9161	Da	
Do Bot Do	Do	- Cyclobarbital No. 04-9175	Do	Do.
Do Heroin No. 04-9162 Do Do Do. Do Hexobarbital No. 04-9163 Do Do. Do. Do. Do Methadone No. 04-9163 Do Do. Do. Do. Do Methadone No. 04-9163 Do Do. Do. Do. Do Methamphetamine No. 04-9163 Do Do. Do. Do. Do Mephobarbital No. 04-9160 Do Do. Do. Do. Do Pentobarbital No. 04-9161 Do Do. Do. Do. Do Phenobarbital No. 04-9181 Do Do. Do. Do. Do Phenobarbital No. 04-9180 Do Do. Do. Do. Do Secobarbital No. 04-9180 Do Do. Do. Do. Do. Do Barb mix-1 Do Do Do. Do. <td></td> <td>- Giuterununde 140, 04-2110</td> <td></td> <td></td>		- Giuterununde 140, 04-2110		
Do Hexobarbital No. 04-9177 Do Do. Do Methadone No. 04-9173 Do Do Do. Do Methadone No. 04-9163 Do Do. Do. Do Morphine No. 04-9160 Do Do. Do. Do Pentobarbital No. 04-9179 Do Do. Do. Do Phenobarbital No. 04-9180 Do Do. Do. Do Secobarbital No. 04-9180 Do Do. Do. Do Secobarbital No. 04-9180 Do Do. Do. Do Barb mix-1 Do Do. Do. Do. Do Amph mix Do Do Do. Do. Do. Do. Do. Do.	Do	Heroin No. 04-9169	Do	
Do Methadone No. 04-9163 Do Do Do Methamphetamine No. 04-9163 Do Do Do Do Methamphetamine No. 04-9163 Do Do Do Do Methamphetamine No. 04-9178 Do Do Do Do Morphine No. 04-9178 Do Do Do Do Pentobarbital No. 04-9179 Do Do Do Do Pentobarbital No. 04-9179 Do Do Do Do Phenobarbital No. 04-9180 Do Do Do Do Secobarbital No. 04-9180 Do Do Do Do Do Secobarbital No. 04-9180 Do	Do	- Hevobarbital No. 04-9177	Do	
Do Methamphetamine No. 04-9168 Do Do Do. Do Mephobarbital No. 04-9178 Do Do. Do. Do Pentobarbital No. 04-9169 Do Do. Do. Do Pentobarbital No. 04-9179 Do Do. Do. Do Pentobarbital No. 04-9179 Do Do. Do. Do Phenobarbital No. 04-9179 Do Do. Do. Do Phenobarbital No. 04-9180 Do Do. Do. Do Secobarbital No. 04-9180 Do Do. Do. Do Secobarbital No. 04-9180 Do Do. Do. Do Barb mix-1 Vial: 1 ml. Aug. 28, 1973 Do Barb mix-2 Do Do. Do. Do Ampho mix Do Do. Do. Do Alk mix Do Do. Do. Do Cannabichromene, No. 04-9220 Ampoule: 1 ml. Nov. 27, 1974 Do Cannabidiol, No. 04-9223 Do	Do	_ Methadone No. 04-9163	Do ano do	
Do Morphine No. 04-9160 Do Do Do Do Pentobarbital No. 04-9179 Do Do<		_ Methamphetamine No. 04-9168	Do	
Do Pentobarbital No. 04-9179 Do Do Do Do Phenobarbital No. 04-9180 Do Do <t< td=""><td></td><td>_ Mephobarbital No. 04-9178</td><td>Do</td><td></td></t<>		_ Mephobarbital No. 04-9178	Do	
Do Phenobarbital No. 04-9181 Do Do Do Phenylmethylbarbituric acid No. Do Do Do 04-9182. Do Do Do Do Do Do Secobarbital No. 04-9180 Do Do Do Do Do Barb mix-1 Vial: 1 ml. Aug. 28, 1973 Do Do Do Do Barb mix-1 Do		Pentobarbitel No. 04-9170	Do	
Do Phenylmethylbarbituric acid No. Do Do. 04-9182. Secobarbital No. 04-9180 Do Do. Do. Do Barb mix-1 Vial: 1 ml. Aug. 28, 1973 Do Barb mix-2 Do Do. Do. Do Amph mix Do Do. Do. Do Alk mix Do Do. Do. Do Cannabichromene, No. 04-9220 Ampoule: 1 ml. Nov. 27, 1974 Do Cannabidiol, No. 04-9223 Do Do. Do. Do. Do Cannabidiol, No. 04-9223 Do Do. Do. Do.	Do	Phenobarbital No. 04-9181	Do	
Do Barb mix-1 Vial: 1 ml. Aug. 28, 1973 Do Barb mix-2 Do Do Do Do Amph mix Do Do Do Do Amph mix Do Do Do Do Cannabichromene, No. 04-9220 Ampoule: 1 ml. Nov. 27, 1974 Do Cannabidiolic Acid, No. 04-9222 Do Do Do Cannabidiolic Acid, No. 04-9223 Do Do		Phenylmethylbarbituric acid No. 04-9182.	Do	
Do Barb mix-1 Vial: 1 ml. Aug. 28, 1973 Do Barb mix-2 Do Do Do Do Amph mix Do Do Do Do Cannabichromene, No. 04-9220 Ampoule: 1 ml. Nov. 27, 1974 Do Cannabidiolic Acid, No. 04-9222 Do Do Do Cannabidiolic Acid, No. 04-9223 Do Do			Do	Do.
Do Amph mix Do Do. Do Alk mix Do Do. Do Cannabichromene, No. 04-9220 Ampoule: 1 ml. Nov. 27, 1974 Do Cannabidiol, No. 04-9221 Do Do. Do Cannabidiol, No. 04-9222 Do Do. Do Cannabidiol, No. 04-9223 Do Do. Do Cannabidioli Acid, No. 04-9223 Do Do.		Barb mix-1	Vial: 1 ml	Aug. 28, 1973
Do Alk mix Do Do. Do Cannabichromene, No. 04-9220 Ampoule: 1 ml. Nov. 27, 1974 Do Cannabidiol, No. 04-9221 Do Do Do. Do Cannabidiol, No. 04-9221 Do Do. Do. Do Cannabidiolic Acid, No. 04-9222 Do Do. Do. Do Cannabidiolic Acid, No. 04-9223 Do Do. Do.	Do	Amph mix	D0	
Do Cannabichromene, No. 04-9220 Ampoule: 1 ml. Nov. 27, 1974 Do Cannabidiol, No. 04-9221 Do Do. Do Cannabidiolic Acid, No. 04-9222 Do Do. Do Cannabigerol, No. 04-9223 Do Do. Do Cannabigerol, No. 04-9223 Do Do. Do Cannabigerol, No. 04-9223 Do Do. Do Cannabigerol, No. 04-9225 Do Do.			Do	
Do Cannabidiol, No. 64-9221 Do Do. Do Cannabidiolic Acid, No. 04-9222 Do Do. Do Cannabidiolic Acid, No. 04-9223 Do Do. Do Cannabidiolic Acid, No. 04-9223 Do Do. Do Cannabigerol, No. 04-9223 Do Do. Do Cannabinol, No. 04-9225 Do Do.		_ Cannabichromene, No. 04-9220	Ampoule: 1 ml.	Nov. 27, 1974
Do Cannabidiolic Acid, No. 04-9222 Do Do. Do	Do	Cannabidiol. No. 04-9221	Do	Do.
Do Do Do.	Do	_ Cannabidiolic Acid, No. 04-9222 _	Do	Do.
Do Cannabinol, No. 04-0226 Do Do		_ Cannabigerol, No. 04-9223		
	Do	Campabinol, No. 04-0225	U0	Do. Do.

Manu	facturer or supplier	Product name and supplier's catalog number	Form of product	Date of application
Do Do		Δ ¹ -THC, No. 04-9227 Δ ⁶ -THC, No. 04-9228	Do	Do. Do.
	Co	Frat benzoyl ecgonine calibrator Frat methadone calibrator	Vial: 1 ml.	Sept. 13, 197
Do	******************	Frat opiate calibrator	Do	Do.
			Do	Do.
Do		Frat barbiturate calibrator	Do	Do. Do.
Do			Bottle: 5 ml.	May 22, 197
Do	**	agent B.	Do	Do.
Do	****	Frat Barbiturate Spin Label Re- agent B.	. –	Do.
Do Do	******	Frat Amphetamine Spin Label Re- agent B.		Do.
_		Frat Cocaine Metabolite Spin Label Reagent B.		Do.
		Emit Opiate Enzyme Reagent B	Do	Do.
20		Emit Methadone Enzyme Reagent B.		Do,
Do		Emit Barbiturate Enzyme Reagent B.	Do	Do.
Do		Emit Amphetamine Enzyme Re- agent B.	Do	Do.
Do		Emit Cocaine Metabolite Enzyme Reagent B.	Do	Do.
Do		Emit Onista Essent Dataset D	D.443	_
		Emit Opiate Enzyme Reagent B. Emit Methadone Enzyme Reagent B.	Bottle: 60 ml.	Po. Do,
Do		Emit Barbiturate Enzyme Reagent B.	Do	Do.
Do	*		Do	Do.
Do		Emit Cocaine Metabolite Enzyme Reagent B.	Do	Do.
Do Do	******	Emit High Calibrator Products of the following sub-	Do Vial: 60 ml	May 5, 1973 May 31, 1973
		stances either alone or in com- bination with one another and not to exceed 20 micrograms per milliliter lyophilized human urine. (1) Amphetamine (2) Benzoyl Ecgonine, (3) Codeine, (4) Ecgonine, (5) 2-Ethilidene- 1,5-dimethyl-3,3-diphenyl-pyrroli- dine, (6) Glutchimide, (7) Meth- adone, (8) Methamphetamine, (9) Methaqualone, (10) Mor- phine, (11) Morphine Glucuro- nide, (12) Pentobarbital, (13) Phenobarbital, (14) Secobarbital.	-	Jady 11, 191
Do		Emit AED-No. 1 Calibrator	Vial: 3 ml., lyophilized	Aug. 27, 1976
Do		Emit AED-No. 2 Calibrator	Do	Do.
		Emit AED-No. 3 Calibrator	Do	Do.
Do		Emit AED-No. 4 Calibrator	Do	Do.
Do		Emit AED-No. 5 Calibrator	Do	Do.
		Emit AED-No. 1 Calibrator	Vial: 10 ml. lyophilized	Do.
		Emit AED-No. 2 Calibrator		
		Fait AFD No. 9 California	Do	Do.
		Emit AED-No. 3 Calibrator	Do	Do.
		Emit AED-No. 4 Calibrator	Do	Do.
		Emit AED-No. 5 Cailbrator	Do	Do.
		Antiepileptic Drug Control	Viai: 10 ml. lyophilized	Do.
Do		Emit Phenobarbital Enzyme Re- agent B.	Vial: 5.5 ml., lyophil- ized.	Do.

Manufacturer or supplier	Product name and supplier's catalog number	Form of product	Date of application
Do	Coulter Tox Opiate Enzyme Re- agent.	Vial: 1 and 2 ml.	Do.
Do	Coulter Tox Methadone Enzyme Reagent.	Do	Do.
Do	Coulter Tox Amphetamine Enzyme Reagent.	Do	Do.
Do	Coulter Tox Barbiturate Enzyme Reagent.	Do	Do. :
Do	Coulter Tox Cocaine Metabolite Enzyme Reagent.	Do	Do.
Do	Emit DAU LOW Calibrator	Vial: 8 ml.	July 29, 1975
Do	Emit DAU MEDIUM Calibrator	Do	Do.
Do	Emit BENNODIAZEPINE ME- TABOLITE Enzyme Resgent B.	Glass bottle: 5.5 ml	Do.
Do	Do	Glass bottle: 50.0 ml	Do.
Taylor Chemicals, Inc	Code 1807D-Zinc Reagent D	Bottle: 2 oz, 4 oz, 8 oz, 16 oz.	Aug. 81, 1978
Do	Special Zinc Reagent catalog No. 1807-D.S.	Bottle: 1 qt. (82 fl oz), 1 pt (16 fl oz), 4 oz (4 fl oz).	Sept. 29, 1975
Fechnam, Inc	Benzoyl Ecgonine-BSA, lot No. 81- 172-A.	Glass vial: 8 ml.	July 21, 1975
Do	Benzoyi Ecgonine-RSA, lot No. 81- 172-B.	Do	Do.

(Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.25; filed Jul 9, 1974, 9:29 am: Unpublished; filed Jun 9, 1977, 8:55 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-12 Rulemaking hearings

Authority: IC 35-48-3-1 Affected: IC 4-22-2; IC 35-48-3-1

Sec. 12. Hearings for rule making. In any case where the Indiana Board of Pharmacy shall hold a hearing on the issuance, amendment, or repeal of rules pursuant to IC 1971, 35-24.1-2 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended, the procedures for such hearing and accompanying proceedings shall be governed generally by the rule making procedures set forth in IC 1971, 4-22-2 as amended, and such procedures, if relating to standards and schedules, be of record in accordance with IC 1971, 35-24.1-2-1 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended. (Indiana Board of Pharmacy; Reg 28, Ch II,Sec 2.31; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-2-13 Purpose of public hearings

Authority:	IC 35-48-3-1
Affected:	IC 35-48-3-1

Sec. 13. Purpose of hearing. Whenever proceedings are initiated pursuant to IC 1971, 35-24.1-2 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended, of this chapter, the Indiana Board of Pharmacy shall hold a hearing of record for the purpose of receiving factual evidence and expert opinion regarding the issues involved in the issuance, amendment or repeal of a rule issuable pursuant to IC 1971, 35-24.1-2 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended.

On the date set for hearing any interested party in person or by attorney shall be afforded an adequate opportunity to participate

in the formulation of the proposed rule or rules through the presentation of facts or argument or the submission of written data or views. All relevant matter presented shall be given full consideration by the Board.

The Board may adopt procedures in addition to those required by this Act *[IC 35-48]* including the holding of conferences and inviting and permitting the submission of suggestions, facts, argument and views of interested persons in advance of the drafting of the proposed rule or rules. *(Indiana Board of Pharmacy; Reg 28, Ch II, Sec 2.32; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)*

856 IAC 2-2-14 Exempt anabolic steroid products

Authority: IC 35-48-2-14 Affected: IC 35-48-2

Sec. 14. The following anabolic steroid containing compounds, mixtures, or preparations have been exempted from this rule and are not controlled substances:

Trade Name	Composition	Company
Androgyn L.A.	Vial; testosterone enathate 90 mg-ml; estradiol valerate 4 mg-ml	Forest Pharmaceuticals St. Louis, MO
Andro-estro 90-4	Vial; testosterone enanthate 90 mg-ml; estradiol valerate 4 mg-ml	Rugby Laboratories Rockville Centra, NY
depANDROGYN	Vial; testosterone cypionate 50 mg-ml; estradiol cypionate 2 mg-ml	Forest Pharmaceuticals St. Louis, MO
DEPO-T.E.	Vial; testosterone cypionate 50 mg-ml; estradiol cypionate 2 mg-ml	Quality Research Pharmaceuticals Carmel, IN
deptestROGEN	Vial; testosterone cypionate 50 mg-ml; estradiol cypionate 2 mg-ml	Martica Pharmaceuticals Phoenix, AZ
Duomone	Vial; testosterone enanthate 90 mg-ml; estradiol valerate 4 mg-ml	Wintec Pharmaceuticals Pacific, MO
DURAtestRIN	Vial; testosterone cypionate 50 mg-ml; estradiol cypionate 2 mg-ml	W.E. Hauck Alpharetta, GA
DUO-SPAN II	Vial; testosterone cypionate 50 mg-ml; Esterified cypionate 2 mg-ml	Primedics Laboratories Gardena, CA
Estratest	Tablet; esterified estrogens 1.25 mg; methyltestosterone 2.5 mg	Solvay Pharmaceuticals Marietta, GA
Estratest HS	Tablet; esterified estrogens 0.625 mg; methyltestosterone 1.25 mg	Solvay Pharmaceuticals Marietta, GA
PAN estra test	Vial; testosterone cypionate 50 mg-ml; estradiol cypionate 2 mg-ml	Pan American Labs Covington, LA
Premarin with methyltestosterone	Tablet; conjugated estrogens 1.25 mg; methyltestosterone 10.0 mg	Ayerst Labs, Inc. New York, NY
Premarin with methyltestosterone	Tablet; conjugated estrogens 0.625 mg; methyltestosterone 5.0 mg	Ayerst Labs, Inc. New York, NY
Test-ESTRO cypionates	Vial; testosterone cypionate 50 mg-ml; estradiol cypionate 2 mg-ml	Rugby Laboratories Rockville Center, NY
Testosterone Cyp 50 estradiol Cyp 2	Vial; testosterone cypionate 50 mg-ml; estradiol cypionate 2 mg-ml	I.D.E Interstate Amityville, NY

Testosterone cypionate-	Vial; testosterone cypionate 50 mg-ml;	Best Generics
estradiol cypionate injection	estradiol cypionate 2 mg-ml	No. Miami Beach, FL
Testosterone cypionate-	Vial; testosterone cypionate 50 mg-ml;	Goldline Labs
estradiol cypionate injection	estradiol cypionate 2 mg-ml	Ft. Lauderdale, FL
Testosterone cypionate-	Vial; testosterone cypionate 50 mg-ml;	Scein Pharmaceuticals
estradiol cypionate injection	estradiol cypionate 2 mg-ml	Port Washington, NY
Testosterone cypionate-	Vial; testosterone cypionate 50 mg-ml;	Steris Labs, Inc.
estradiol cypionate injection	estradiol cypionate 2 mg-ml	Phoenix, AZ
Testosterone enanthate-	Vial; testosterone enanthate 90 mg-ml;	Steris Labs, Inc.
estradiol valerate injection	estradiol valerate 4 mg-ml	Phoenix, AZ

(Indiana Board of Pharmacy; 856 IAC 2-2-14; filed May 31, 1994, 5:00 p.m.: 17 IR 2337; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

Rule 3. Registration Information–Special Instructions

856 IAC 2-3-1 Registration information furnished upon request

Authority: IC 35-48-3-1 Affected: IC 35-48-3-1

Sec. 1. Information; special instructions. Information regarding procedures under these rules and instructions supplementing these rules will be furnished upon request by writing to the Indiana State Board of Pharmacy. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.02; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-2 Persons required to register

Authority: IC 35-48-3-1 Affected: IC 35-48-3-3

Sec. 2. Persons required to register. Every person who manufactures, distributes, or dispenses any controlled substance or who proposes to engage in the manufacture, distribution, or dispensing of any controlled substance shall obtain annually a registration unless exempted by law or pursuant to sections 3.14-3.17 [856 IAC 2-3-5 - 856 IAC 2-3-8] of this chapter. Only persons actually engaged in such activities are required to obtain registration; related or affiliated persons who are not engaged in such activities are not required to be registered. (For example, a stockholder or parent corporation of a corporation manufacturing controlled substances is not required to obtain a registration).

As soon after the effective date of these rules as is practicable, the Board shall issue a provisional certificate to all persons when in possession of a valid State of Indiana or Federal certificate of registration authorizing such persons to manufacture, distribute, dispense, prescribe or possess controlled substances. The provisional certificates shall be valid until the Board shall declare that applications for annual renewals shall begin and until such applications have been acted upon by the Board. During the first renewal period, when it is instituted, applications shall be required from all prospective registrants in alphabetically ordered increments according to a schedule to be adopted by the Board. (*Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.11; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA*)

856 IAC 2-3-3 Independent activities; separate registration required; exceptions

Authority: IC 35-48-3-1 Affected: IC 35-48-3-3; IC 35-48-3-4

Sec. 3. Separate registration for independent activities. (a) The following groups of activities are deemed to be independent

of each other:

(1) Manufacturing controlled substances;

(2) Distributing controlled substances;

(3) Dispensing controlled substances listed in Schedules II through V [856 IAC 2-2-3 - 856 IAC 2-2-6];

(4) Conducting research (other than research described in sub-paragraph (6) of this paragraph) with controlled substances listed in Schedules II through V [856 IAC 2-2-3 - 856 IAC 2-2-6];

(5) Conducting instructional activities with controlled substances listed in Schedules II through V [856 IAC 2-2-3 - 856 IAC 2-2-61:

(6) Conducting research with narcotic drugs listed in Schedules II through V /856 IAC 2-2-3 - 856 IAC 2-2-6] for the purpose of continuing the dependence on such drugs of a narcotic drug dependent person in the course of conducting an authorized clinical investigation in the development of a narcotic addict rehabilitation program pursuant to a Notice of Claimed Investigational Exemption for a New Drug approved by the Food and Drug Administration;

(7) Conducting research and instructional activities with controlled substances listed in Schedule I [856 IAC 2-2-2]; and (8) Conducting chemical analysis with controlled substances listed in any Schedule.

(b) Every person who engages in more than one group of independent activities shall obtain a separate registration for each group of activities except as provided in this paragraph. Any person, when registered to engage in the group of activities described in each subparagraph in this paragraph, shall be authorized to engage in the coincident activities described in that subparagraph without obtaining a registration to engage in such coincident activities, provided that, unless specifically exempted, he complies with all requirements and duties prescribed by law for persons registered to engage in such coincident activities:

(1) A person registered to manufacture any controlled substance or basic class of controlled substance shall be authorized to distribute that substance or class, but no other substance or class which he is not registered to manufacture;

(2) A person registered to manufacture any controlled substance listed in Schedules II through V /856 IAC 2-2-3 - 856 IAC 2-2-6] shall be authorized to conduct chemical analysis and pre-clinical research (including quality control analysis) with narcotic and non-narcotic controlled substances listed in those schedules in which he is authorized to manufacture;

(3) A person registered to conduct research with a basic class of controlled substance listed in Schedule I [856 IAC 2-2-2] shall be authorized to manufacture such class if and to the extent that such manufacture is set forth in a research protocol federally approved by the Drug Enforcement Administration and to distribute such class to other persons registered or authorized to conduct research with such class or registered or authorized to conduct chemical analysis with controlled substances;

(4) A person registered or authorized to conduct chemical analysis with controlled substances shall be authorized to manufacture such substances for analytical or instructional purposes, to distribute such substances to other persons registered or authorized to conduct chemical analysis or instructional activities or research with such substances and to persons exempted from registration pursuant to section 3.16 [856 IAC 2-3-7], and to conduct instructional activities with controlled substances; and

(5) A person registered or authorized to conduct research (other than research described in paragraph (a)(6) of this section) with controlled substances listed in Schedules II through V /856 IAC 2-2-3 - 856 IAC 2-2-6] shall be authorized to conduct chemical analysis with controlled substances listed in those schedules in which he is authorized to conduct research, to manufacture such substances if and to the extent that such manufacture is set forth in a statement filed with the application for registration, to distribute such substances to other persons registered or authorized to conduct chemical analysis, exempted from registration pursuant to Section 3.16 /856 IAC 2-3-7], and to conduct instructional activities with controlled substances; (6) A person registered to dispense controlled substances listed in Schedules II through V [856 IAC 2-2-3 – 856 IAC 2-2-6] shall be authorized to conduct research (other than research described in paragraph (a) (6) of this section) and to conduct instructional activities with those substances.

(7) A person registered as a manufacturer shall be authorized to conduct one, all or several of the activities and coincident activities enumerated and described in paragraphs (b)(1), (b)(2), (b)(3), (b)(4), and (b)(5) under a single registration if set forth in his application and pertaining to those controlled substances or schedules as set forth in his application. (For example, a manufacturer under a single registration may perform all or any of the following activities, by way of illustration and not limitation; (a) manufacture and distribute any controlled substance or basic class, (b) chemical analysis, (c) Schedule I /856 IAC 2-2-2] research pursuant to a federally approved protocol, (d) Schedule II through V [856 IAC 2-2-3 – 856 IAC 2-2-6] research, and (e) instructional activity if set forth in his application and for those controlled substances or schedules as set forth for each activity.

(c) A single registration to engage in any group of independent activities may include one or more controlled substances listed

in the schedules authorized in that group of independent activities. A person registered to conduct research with controlled substances listed in Schedule I [856 IAC 2-2-2] may conduct research with any substance listed in Schedule I [856 IAC 2-2-2] for which he has filed and had approved a research protocol, by the Federal Drug Enforcement Administration. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.12; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-4 Separate registrations for separate locations; exceptions

Authority: IC 35-48-3-1 Affected: IC 35-48

Sec. 4. Separate registrations for separate locations. (a) A separate registration is required for each principal place of business or professional practice at one general physical location where controlled substances are manufactured, distributed, dispensed, or possessed by a person.

(b) The following locations shall be deemed not to be places where controlled substances are manufactured, distributed, or dispensed:

(1) A warehouse where controlled substances are stored by or on behalf of a registered person, unless such substances are distributed directly from such warehouse to registered locations other than the registered location from which the substances were delivered or to persons not required to register by virtue of IC 1971, 35-24.1-3-2(c)(2) [Repealed by P.L.26-1977, SECTION 25. Compare IC 35-48-3-3.] as amended.

(2) An office used by agents of a registrant where sales of controlled substances are solicited, made, or supervised but which neither contains such substances (other than substances for display purposes or lawful distribution as samples only) nor serves as a distribution point for filling sales orders; and

(3) An office used by a practitioner (who is registered at another location) where controlled substances are prescribed but neither administered nor otherwise dispensed as a regular part of the professional practice of the practitioner at such office, and where no supplies of controlled substances are maintained.

(c) The requirement of registration is waived for ambulances as defined by IC 16-1-39-2 [IC 16-1 was repealed by P.L.2-1993, SECTION 209, effective July 1, 1993.] and 836 IAC 1-1-1 operated by an ambulance service provider also defined at 836 IAC 1-1-1 which holds certification as a provider organization as this term is defined in IC 16-1-40 [IC 16-1 was repealed by P.L.2-1993, SECTION 209, effective July 1, 1993.] from the Indiana Emergency Medical Services Commission, providing that the pharmacies of the supervising or sponsoring hospitals hold a valid Indiana Board of Pharmacy permit and valid Indiana and Federal Controlled Substances Registration. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.13; filed Jul 9, 1974, 9:29 am: unpublished; filed Feb 11, 1981, 9:05 am: 4 IR 377; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-5 Exemption of agents or employees; affiliated practitioners; paramedics

Authority: IC 35-48-3-1 Affected: IC 35-48-3-3

Sec. 5. Exemption of agents and employees; affiliated practitioners. (a) The requirement of registration is waived for any agent or employee of a person who is registered to engage in any group of independent activities, if such agent or employee is acting in the usual course of his business or employment.

(b) An individual practitioner (other than an intern, resident, or foreign-trained physician) who is an agent or employee of another practitioner registered to dispense controlled substances may, when acting in the usual course of his employment, administer, and dispense (other than by issuance of prescription) controlled substances if and to the extent that such individual practitioner is authorized or permitted to do so by the jurisdiction in which he practices, under the registration of the employer or principal practitioner in lieu of being registered himself. (For example, a pharmacist employed by a pharmacy need not be registered individually to fill a prescription for controlled substances in a pharmacy if so registered).

(c) An individual practitioner who is an intern, resident, or foreign-trained physician may dispense, administer, and prescribe controlled substances under the registration of the hospital or other institution which is registered and by whom he is employed in lieu of being registered himself, provided that:

(1) Such dispensing or prescribing is done in the usual course of his professional practice;

(2) Such individual practitioner is authorized or permitted to do so by the jurisdiction in which he is practicing;

(3) The hospital or other institution by whom he is employed has determined that the individual practitioner is so permitted to dispense, administer, or prescribe drugs by the jurisdiction;

(4) Such individual practitioner is acting only within the scope of his employment with the hospital or institution;

(5) The hospital or other institution authorizes the intern, resident, or foreign-trained physician to dispense or prescribe under the hospital registration and designates a specific internal code number for each intern, resident, or foreign-trained physician so authorized. The code number shall consist of numbers, letters, or a combination thereof and shall be a suffix to the institution's DEA registration number, preceded by a hyphen (e.g., AP 0123456-10 or AP 0123456-A12); and

(6) A current list of internal codes and the corresponding individual practitioner is kept by the hospital or other institution and is made available to the public upon request for the purpose of verifying the authority of the prescribing individual practitioner.(d) The requirement of registration is waived for advanced emergency medical technicians and emergency parametics as

(d) The requirement of registration is warved for advanced energency medical technicians and energency parametrics as described in IC 16-1-40 [IC 16-1 was repealed by P.L.2-1993, SECTION 209, effective July 1, 1993.] and 836 IAC 2-1-1 insofar as they administer controlled substances within the applicable requirements and standards of IC 16-1-40 [IC 16-1 was repealed by P.L.2-1993, SECTION 209, effective July 1, 1993.] as well as the rules and regulations promulgated thereunder by the Indiana Emergency Medical Services Commission. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.14; filed Jul 9, 1974, 9:29 am: unpublished; filed Feb 11, 1981, 9:05 am: 4 IR 378; errata, 4 IR 536; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-6 Exemption of military or public health service personnel

Authority: IC 35-48-3-1 Affected: IC 35-48-3-3

Sec. 6. Exemption of certain military and other personnel. (a) The requirement of registration is waived for any official of the U.S. Army, Navy, Marine Corps, Air Force, Coast Guard, or Public Health Service who is authorized to prescribe, dispense, or administer, but not to procure or purchase, controlled substances in the course of his official duties. Such officials shall follow procedures set forth in Part 6 *[856 IAC 2-6]* of this chapter regarding prescriptions, but shall state the branch of service or agency (e.g., "U.S. Army" or "Public Health Service") and the service identification number of the issuing official in lieu of the registration number required on prescription forms. The service identification number for a Public Health Service employee is his Social Security identification number.

(b) If any official exempted by this section also engages as a private individual in any activity or group of activities for which registration is required, such official shall obtain a registration for such private activities. (Indiana Board of Pharmacy; Reg 28,Ch III,Sec 3.15; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-7 Exemption of law enforcement officers; registration of law enforcement laboratories

Authority: IC 35-48-3-1 Affected: IC 35-48-3-3

Sec. 7. Exemption of law enforcement officials. (a) The requirement of registration is waived for the following persons in the circumstances described in this section:

(1) Any officer or employee of the Drug Enforcement Administration, any officer of the U.S. Bureau of Customs, any officer or employee of the United States Food and Drug Administration, and any other Federal officer who is lawfully engaged in the enforcement of any Federal law relating to controlled substances, drugs or customs, and is duly authorized to possess controlled substances in the course of his official duties; and

(2) Any officer or employee of any State, or any political subdivision or agency thereof, who is engaged in the enforcement of any State or local law relating to controlled substances and is duly authorized to possess controlled substances in the course of his official duties.

(b) Any official exempted by this section may, when acting in the course of his official duties, possess any controlled substance and distribute any such substance to any other official who is also exempted by this section and acting in the course of his official duties.

(c) Any official exempted by this section may procure any controlled substance in the course of an inspection, in accordance

with IC 1971, 35-24.1-5-2 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended, or in the course of any criminal investigation involving the person from whom the substance was procured.

(d) In order to enable law enforcement agency laboratories to obtain and transfer controlled substances for use as standards in chemical analysis, such laboratories must obtain annually a registration to conduct chemical analysis. Such laboratories shall be exempted from payment of a fee for registration. Laboratory personnel, when acting in the scope of their official duties, are deemed to be officials exempted by this section and within the activity described in IC 1971, 35-24.1-5-6(c) [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended. For the purpose of this paragraph, laboratory activities shall not include field or other preliminary chemical tests by officials exempted by this section. (Indiana Board of Pharmacy; Reg 28, Ch III,Sec 3.16; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-8 Exemption of civil defense officials

Authority: IC 35-48-3-1 Affected: IC 35-48-3-3

Sec. 8. Exemption of civil defense officials. (a) The requirement of registration is waived for any offical of a civil defense or disaster relief organization who, in the course of his official duties, is authorized to:

(1) Maintain, and distribute for such maintenance, controlled substances held for emergency use; or

(2) Procure controlled substances for the purpose of maintaining supplies for emergency use, provided that all of such procurement is from the U.S. General Services Administration and in accordance with the rules of the U.S. Office of Emergency Preparedness.

(b) The requirement of registration is waived for any official of a civil defense or disaster relief organization during a state of emergency or disaster within his jurisdiction proclaimed by the President or by a concurrent resolution of the Congress, which official, in the course of his official duties during such emergency or disaster, is authorized to:

(1) Dispense controlled substances; or

(2) Procure or distribute controlled substances, provided that all such procurement is on a special "Civil Defense Emergency Order Form," as described in this section.

(c) Civil Defense Emergency Order Forms shall be furnished by the U.S. Office of Emergency Preparedness and will contain the name of the civil defense or disaster relief organization. Such forms may be used and are valid only during a state of emergency or disaster proclaimed by the President or by a concurrent resolution of the Congress for the area in which the organization using such forms has civil defense or disaster relief jurisdiction, who shall state his position and the nature and legal designation of the emergency or disaster. Such forms may be filled by any person registered under the Act *[IC 35-48]*. The Organization shall, upon the execution of a Civil Defense Emergency Order Form, be deemed to be registered under the Act *[IC 35-48]* for purposes or recordkeeping pursuant to Part 4 *[856 IAC 2-4]*. *(Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.17; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)*

856 IAC 2-3-9 Registration fees

Authority: IC 25-26-13-4

Affected: IC 25-26-13-4

Sec. 9. (a) For each registration or reregistration to manufacture controlled substances, the registrant shall pay a fee of one hundred dollars (\$100).

(b) For each registration or reregistration to distribute controlled substances, the registrant shall pay a fee of one hundred dollars (\$100).

(c) For each registration or reregistration to dispense or to conduct research or instructional activities with controlled substances listed in 856 IAC 2-2-3 through 856 IAC 2-2-6, the registrant shall pay a fee of one hundred dollars (\$100).

(d) For each registration or reregistration to conduct research or instructional activities with controlled substances listed in 856 IAC 2-2-2, the registrant shall pay a fee of one hundred dollars (\$100).

(e) For each registration or reregistration to conduct chemical analysis with controlled substances listed in any schedule, the registrant shall pay a fee of one hundred dollars (\$100).

(f) For each registration or reregistration for a practitioner seeking to prescribe, administer, or dispense controlled substances, the registrant shall pay a fee of sixty dollars (\$60). (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.21; filed Jul 9, 1974, 9:29 a.m.: Unpublished; filed Jul 8, 1981, 9:00 a.m.: 4 IR 1499; filed Jul 20, 1984, 10:00 a.m.: 7 IR 2379; filed Aug 21, 1986, 10:30 a.m.: 10 IR 63; filed Jun 6, 1996, 9:00 a.m.: 19 IR 3106; readopted filed Oct 17, 2001, 3:25 p.m.: 25 IR 940; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-10 Time and method of payment; refund (Repealed)

Sec. 10. (Repealed by Indiana Board of Pharmacy; filed Nov 30, 2001, 11:00 a.m.: 25 IR 1344)

856 IAC 2-3-11 Persons exempt from fee

Authority: IC 35-48-3-1 Affected: IC 35-48-3-1

Sec. 11. Persons exempt from fee. (a) The Indiana State Board of Pharmacy shall exempt from payment of a fee for registration or re-registration the following persons:

(1) Any official or agency of the U.S. Army, Navy, Marine Corps, Air Force, Coast Guard, Veterans' Administration or Public Health Service who or which is authorized to procure or purchase controlled substances for official use; and

(2) Any official, employee, or other civil officer or agency of the United States, or any State, or any political subdivision or agency thereof, who or which is authorized to purchase controlled substances, to obtain such substances from official stocks, to dispense or administer such substances, to conduct research, instructional activities, or chemical analysis with such substances, or any combination thereof, in the course of his or its official duties or employment.

(b) In order to claim exemption from payment of a registration or re-registration fee, the registrant shall have completed the certification on the appropriate form, wherein the registrant's superior (if an individual) or officer (if an agency) certifies to the status and address of the registrant and to the authority of the registrant to acquire, possess, or handle controlled substances.

(c) Exemption from payment of a registration or re-registration fee does not relieve the registrant of any other requirements or duties prescribed by law. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.23; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-12 Time for registration or re-registration application

Authority: IC 35-48-3-1 Affected: IC 35-48-3-5

Sec. 12. Time for application for registration; expiration date. (a) Any person who is required to be registered and who is not so registered may apply for registration at any time. No person required to be registered shall engage in any activity for which registration is required until the application for registration is granted by the Indiana Board of Pharmacy.

(b) Any person who is registered may apply to be re-registered not more than 60 days, before the expiration date his registration. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.31; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-13 Application forms; reregistration forms

Authority: IC 35-48-3-1 Affected: IC 35-48-3-5

Sec. 13. (a) If any person is required to be registered, and is not so registered and is applying for registration, the following apply:

(1) To manufacture and perform other coincident activities (see 856 IAC 2-3-3(b)(7) [section 3(b)(7) of this rule]) with controlled substances, he or she shall apply on Form CSR-1A.

(2) To dispense, or to conduct research (other than research described in 856 IAC 2-3-3(a)(6) [section 3(a)(6) of this rule]) with, or to conduct instructional activities with, controlled substances listed in Schedules II through V under 856 IAC 2-2-3 through 856 IAC 2-2-6, he or she shall apply on Form CSR-1.

(3) To conduct research with narcotic drugs listed in Schedules II through V under 856 IAC 2-2-3 through 856 IAC 2-2-6, as described in 856 IAC 2-3-3(a)(6) [section 3(a)(6) of this rule], he or she shall apply on Form CSR-1.

(4) To conduct research with controlled substances listed in Schedule I under 856 IAC 2-2-2, he or she shall apply on Form CSR-1 in accordance with an approved Schedule I under 856 IAC 2-2-2 research protocol. Such protocol shall be subject to inspection by the Indiana board of pharmacy.

(5) To conduct instructional activities with controlled substances listed in Schedule I under 856 IAC 2-2-2, he or she shall apply as a researcher on Form CSR-1 with two (2) copies of a statement describing the nature, extent, and duration of such instructional activities attached to the form.

(6) To conduct chemical analysis with controlled substances listed in any schedule, he or she shall apply on Form CSR-1.

(7) To distribute controlled substances, he or she shall apply on Form CSR-1.

(b) If any person is registered and is applying for reregistration, the following apply:

(1) To manufacture and perform other coincident activities (see 856 IAC 2-3-3(b)(7)[section 3(b)(7) of this rule]), with controlled substances, he or she shall apply on Form CSRII-A.

(2) To dispense, or to conduct research (other than research described in 856 IAC 2-3-3(a)(6)[section 3(a)(6) of this rule]) with, or to conduct instructional activities with, controlled substances listed in Schedules II through V under 856 IAC 2-2-3 through 856 IAC 2-2-6, he or she shall apply on Form CSR-II.

(3) To conduct research with narcotic drugs listed in Schedules II through V under 856 IAC 2-2-3 through 856 IAC 2-2-6, as described in 856 IAC 2-3-3(a)(6) [section 3(a)(6) of this rule], he or she shall apply on Form CSR-II.

(4) To continue to conduct research with controlled substances listed in Schedule I under 856 IAC 2-2-2 under one (1) or more approved research protocols, by the Drug Enforcement Administration, he or she shall apply on Form CSR-II.

(5) To continue to conduct instructional activities with controlled substances listed in Schedule I under 856 IAC 2-2-2 under one (1) or more approved instructional statements, he or she shall apply as a researcher on Form CSR-II.

(6) To conduct chemical analysis with controlled substances listed in any schedule, he or she shall apply on Form CSR-II.

(7) To distribute controlled substances, he or she shall apply on Form CSR-II.

(c) Applications for registration may be obtained by writing to the controlled substance division of the Indiana board of pharmacy. Applications for reregistration will be mailed, as applicable, to each registered person approximately sixty (60) days before the expiration date of his or her registration; if any registered person does not receive such forms within forty-five (45) days before the expiration date of his or her registration, he or she must promptly give notice of such fact and request such forms by writing to the controlled substance division of the Indiana board of pharmacy.

(d) Each application for registration to handle any basic class of controlled substance listed in Schedule I under 856 IAC 2-2-2 (except to conduct chemical analysis with such classes), and each application for registration to manufacture a basic class of controlled substance listed in Schedule II under 856 IAC 2-2-3, or to conduct research with any narcotic controlled substance listed in Schedule II under 856 IAC 2-2-3, or to conduct research with any narcotic controlled substance listed in Schedule II under 856 IAC 2-2-3, or to conduct research with any narcotic controlled substance listed in Schedule II under 856 IAC 2-2-3, or to conduct research with any narcotic controlled substance listed in Schedule II under 856 IAC 2-2-3, or to conduct research with any narcotic controlled substance listed in Schedule II under 856 IAC 2-2-3, or to conduct research with any narcotic controlled substance listed in Schedule II under 856 IAC 2-2-3, or to conduct research with any narcotic controlled substance listed in Schedule II under 856 IAC 2-2-3, or to conduct research with any narcotic controlled substance listed in Schedule II under 856 IAC 2-2-3, shall include the controlled substances code number, as set forth in Part I [856 IAC 2-1], for each basic class or substance to be covered by such registration.

(e) Each application shall include all information called for on the form unless the item is not applicable, in which case this fact shall be indicated.

(f) Each application, attachment, or other document filed as part of an application shall be signed by:

(1) the applicant, if an individual;

(2) a partner of the applicant, if a partnership; or

(3) an officer or authorized representative of the applicant, if a corporation, corporate division, association, trust, or other entity.

(Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.32; filed Jul 9, 1974, 9:29 a.m.: Unpublished; readopted filed Nov 30, 2001, 11:00 a.m.: 25 IR 1342; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-14 Filing of application; joint filing (Repealed)

Sec. 14. (Repealed by Indiana Board of Pharmacy; filed Nov 30, 2001, 11:00 a.m.: 25 IR 1344)

856 IAC 2-3-15 Acceptance for filing; defective applications; requests for additional information (Repealed)

Sec. 15. (Repealed by Indiana Board of Pharmacy; filed Nov 30, 2001, 11:00 a.m.: 25 IR 1344)

856 IAC 2-3-16 Additional information; failure to supply

Authority: IC 35-48-3-1 Affected: IC 35-48-3-5

Sec. 16. Additional information. The Indiana State Board of Pharmacy may require an applicant to submit such documents or written statements of fact relevant to the application as the Board deems necessary and as provided by IC 1971, 35-24.1-3-3(a) [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended, to determine whether the application should be granted. The failure of the applicant to provide such documents or statements within a reasonable period of time after being requested to do so shall be deemed to be a waiver by the applicant of an opportunity to present such documents or facts for consideration by the Indiana State Board of Pharmacy in granting or denying the application. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.35; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-17 Amendment or withdrawal of application

Authority:	IC 35-48-3-1
Affected:	IC 35-48-3-5

Sec. 17. Amendments to and withdrawal of applications. An application may be amended or withdrawn without permission of the Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.36; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-18 Inspection and review of application by board

Authority: IC 35-48-3-1 Affected: IC 35-48-3-5

Sec. 18. Administrative review generally. The Indiana Board of Pharmacy may inspect, or cause to be inspected, the establishment of an applicant or registrant, pursuant to IC 1971, 35-24.1-5-2 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended. The Indiana Board of Pharmacy shall review the application for registration and other information regarding an applicant in order to determine whether the applicable standards of IC 1971, 35-24.1-3 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended. Sec IC 35-48.] as amended of Pharmacy shall review the application for registration and other information regarding an applicant in order to determine whether the applicable standards of IC 1971, 35-24.1-3 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended, have been met by the applicant. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.41; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-19 Certificate of registration; denial of registration

Authority: IC 35-48-3-1 Affected: IC 35-48-3-5

Sec. 19. Certificate of registration; denial of registration. (a) The Indiana State Board of Pharmacy shall issue a Certificate of Registration Form CSR-3 to an applicant if the issuance of registration or re-registration is required under the applicable provisions of IC 1971, 35-24.1-3 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended. In the event that the issuance of registration or re-registration is not required, the Indiana State Board of Pharmacy may deny the application. Before denying any application, the Indiana State Board of Pharmacy shall issue an order to show cause pursuant to Section 3.46 [856 IAC 2-3-23] and, if requested by the applicant, shall hold a hearing on the application pursuant to Section 3.51, through Section 3.53 [856 IAC 2-3-24 – 856 IAC 2-3-26].

(b) The Certificate of Registration shall contain the name, address, and registration number of the registrant, the activity authorized by the registration, the schedules and/or Controlled Substances Code Number (as set forth in Part 2 [856 IAC 2-2] of this Act) of the controlled substances which the registrant is authorized to handle, the amount of fee paid (or exemption), and the expiration date of the registration. The registrant shall maintain the Certificate of Registration in a readily retrievable manner and shall permit inspection of the Certificate of Registration and shall permit inspection of the certificate by any official, agent, or employee of the Board or any agency engaged in enforcement of laws relating to controlled substances. (Indiana Board of Pharmacy;

Reg 28, *Ch III*, *Sec* 3.42; *filed Jul* 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-20 Suspension or revocation of registration

Authority:	IC 35-48-3-1
Affected:	IC 35-48-3-5

Sec. 20. Suspension or revocation of registration. (a) The Indiana Board of Pharmacy may suspend any registration pursuant to IC 1971, 35-24.1-3-4 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended, for any period of time he determines.

(b) The Indiana Board of Pharmacy may revoke any registration pursuant to IC 1971, 35-24.1-3-4 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended.

(c) Before revoking or suspending any registration, the Indiana Board of Pharmacy shall issue an order to show cause, such order shall be sent by certified mail to address of the registrant, advising registrant of his rights to a hearing, Form CSR-4, pursuant to section 3.46 *[856 IAC 2-3-23]*. Notwithstanding the requirements of this section, however, the Indiana Board of Pharmacy may suspend any registration pending a final order pursuant to section 3.44 *[856 IAC 2-3-21]*.

(d) Upon service of the final order of the Indiana Board of Pharmacy following a hearing or waiver thereof suspending or revoking registration, the registrant shall immediately deliver his Certificate of Registration to the Indiana Board of Pharmacy. Also, upon service of the final order of the Indiana Board of Pharmacy suspending or revoking registration, the registrant shall, as instructed by the Indiana Board of Pharmacy:

(1) Deliver all controlled substances in his possession to the Indiana Board of Pharmacy or to authorized agents of the Indiana Board of Pharmacy; or

(2) Place all controlled substances in his possession under seal as described in IC 1971, 35-24.1-3-4(c) [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended.

(e) In the event that revocation or suspension is limited to a particular controlled substance or substances, the registrant shall be given a new Certificate of Registration for all substances not affected by such revocation or suspension; no fee shall be required to be paid for the new Certificate of Registration. The registrant shall deliver the old Certificate of Registration to the Indiana Board of Pharmacy. Also, the registrant shall, as instructed by the Indiana Board of Pharmacy:

(1) Deliver to the Indiana Board of Pharmacy or to authorized agents of the Indiana Board of Pharmacy all of the particular controlled substance or substances affected by the revocation or suspension which are in his possession; or

(2) Place all of such substances under seal as described in IC 1971, 35-24.1-3-4(c) [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended.

(Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.43; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-21 Suspension pending final order

Authority: IC 35-48-3-1 Affected: IC 35-48-3-5

Sec. 21. Suspension of registration pending final order. (a) The Indiana Board of Pharmacy may suspend any registration simultaneously with or at any time subsequent to the service upon the registrant of an order to show cause why such registration should not be revoked or suspended, in any case where it finds that there is an imminent danger to the public health or safety. If the Indiana Board of Pharmacy so suspends, it shall serve with the order to show cause pursuant to section 3.46 [856 IAC 2-3-23] an order of immediate suspension which shall contain a statement of its findings regarding the danger to public health or safety.

(b) Upon service of the order of immediate suspension, the registrant shall promptly return his Certificate of Registration to the Indiana Board of Pharmacy. Also, upon service of the order of the Indiana Board of Pharmacy immediately suspending registration, the registrant shall, as instructed by the Indiana Board of Pharmacy:

(1) Deliver all affected controlled substances in his possession to the Indiana Board of Pharmacy or to authorized agents of the Indiana Board of Pharmacy; or

(2) Place all of such substances under seal as described in IC 1971, 35-24.1-3-4(c) [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended.

(c) Any suspension shall continue in effect until the conclusion of all preceedings upon the revocation or suspension, including any judicial review thereof, unless sooner withdrawn by the Indiana Board of Pharmacy or dissolved by a court of competent jurisdiction. Any registrant whose registration is suspended under this section may request a hearing on the revocation or suspension of his registration at a time earlier than specified in the order to show cause pursuant to section 3.46 [856 IAC 2-3-23], which request shall be granted by the Indiana Board of Pharmacy who shall fix a date for such hearing as early as reasonably possible. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.44; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-22 Extension of registration pending re-registration order

Authority: IC 35-48-3-1 Affected: IC 35-48-3-5

Sec. 22. Extension of registration pending final order. In the event that an applicant for re-registration (who is doing business under a registration previously granted and not revoked or suspended) has applied for re-registration at least 30 days before the date on which the existing registration is due to expire, the existing registration of the applicant shall automatically be extended and continue in effect until the date on which the Indiana Board of Pharmacy so issues its final order. The Indiana Board of Pharmacy may extend any other existing registration under the circumstances contemplated in this section even though the registrant failed to apply for re-registration at least 30 days before expiration of the existing registration, with or without request by the registrant, if the Indiana Board of Pharmacy is not inconsistent with the public health and safety. (Indiana Board of Pharmacy; Reg 28, Ch III,Sec 3.45; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-23 Order to show cause

Authority: IC 35-48-3-1 Affected: IC 35-48-3-5

Sec. 23. Order to show cause. (a) If, upon examination of the application for registration from any applicant and other information regarding the applicant, the Indiana Board of Pharmacy is unable to make the determinations required by the applicable provisions of IC 1971, 35-24.1-3-3 *[Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.]*, as amended, to register the applicant, the Indiana Board of Pharmacy shall serve upon the applicant an order to show cause why the registration should not be denied.

(b) If, upon information regarding any registrant, the Indiana Board of Pharmacy determines that the registration of such registrant is subject to suspension or revocation pursuant to IC 1971, 35-24.1-3-4 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.], as amended, the Indiana Board of Pharmacy shall serve upon the registrant an order to show cause why a registration should not be revoked or suspended.

(c) The order to show cause shall call upon the applicant or registrant to appear before the Indiana Board of Pharmacy at a time and place stated in the order, which shall not be less than 30 days after the date of receipt of the order. The order to show cause shall also contain a statement of the legal basis for such hearing and for the denial, revocation, or suspension of registration and a summary of the matters of fact and law asserted.

(d) The Indiana Board of Pharmacy shall hold a hearing at the time and place stated in the order, pursuant to section 3.51 [856 IAC 2-3-24].

(e) When authorized by the section 3.51 [856 IAC 2-3-24] any agent of the Indiana Board of Pharmacy may serve the order to show cause. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.46; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-24 Evidentiary hearing

Authority: IC 35-48-3-1 Affected: IC 35-48-3-6

Sec. 24. The controlled substances advisory committee shall hold a hearing for the purpose of receiving factual evidence regarding the issues involved in the denial, revocation, or suspension of any registration. (Indiana Board of Pharmacy; Reg 28, Ch

III,Sec 3.51; filed Jul 9, 1974, 9:29 a.m.: Unpublished; readopted filed Nov 30, 2001, 11:00 a.m.: 25 IR 1343; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-25 Hearing procedures

Authority: IC 35-48-3-1 Affected: IC 4-21.5; IC 35-48-3-6

Sec. 25. Hearing for granting, denial, revocation, or suspension of application. (a) In any case where the advisory committee shall hold a hearing on any registration or application therefor, the procedures for such hearing shall be governed generally by the adjudication procedures of IC 1971, 4-22-1-1 to 4-22-1-30 *[Repealed by P.L.18-1986, SECTION 2. See IC 4-21.5.]*, as amended, and by sections 3.52-3.53 *[this section and 856 IAC 2-3-26]*.

(b) Any hearing under this part shall be independent of, and not in lieu of, criminal prosecutions or other proceedings under the Act *[IC 35-48]* or any other law of this State.

(c) At any such hearing the advisory committee shall designate one of its members as presiding officer.

(d) At any such hearing a quorum of the advisory committee consisting of a majority of its membership shall hear the evidence and the disputed issues of law and they shall after the conclusion of the hearing, prepare for the Board recommended findings, facts, and conclusions of law.

(e) The committee's recommended findings and facts and conclusions of law shall be acted on by the Board in the manner required by IC 1971, 4-22-1 [Repealed by P.L.18-1986, SECTION 2. See IC 4-21.5.]. (Indiana Board of Pharmacy; Reg 28,Ch III,Sec 3.52; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-26 Modification or waiver of rules

Authority: IC 35-48-3-1 Affected: IC 35-48-3-6

Sec. 26. Waiver or modification of rules. The presiding officer at the advisory committee hearings or of the Indiana Board of Pharmacy (with respect to matters pending before him) may modify or waive any rule in this part by notice in advance of the hearing, if he determines than no party in the hearing will be unduly prejudiced and the ends of justice will thereby be served and if all parties consent. Such notice of modification or waiver shall be made a part of the record of the hearing. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.53; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-27 Modification of registration

Authority: IC 35-48-3-1 Affected: IC 35-48-3-6

Sec. 27. Modification in registration. Any registrant may apply to modify his registration to authorize the handling of additional controlled substances or to change his name or address by submitting a letter of request to the Indiana Board of Pharmacy. The letter shall contain the registrant's name, address, registration number, and the substances and/or schedule to be added to his registration or the name or address and shall be signed by the same person who signed the most recent application for registration or reregistration. If the registrant is seeking to handle additional controlled substances listed in Schedule I *[856 IAC 2-2-2]* for the purpose of research or instructional activities, a Federally approved research protocol describing each research project involving the additional substances shall be subject to inspection by the Indiana Board of Pharmacy or he shall attach two copies of a statement describing the nature, extent, and duration of such instructional activities, as appropriate. No fee shall be required to be paid for the modification. The request for modification shall be handled in the same manner as an application for registration.

If the modification of registration is approved, the Indiana Board of Pharmacy shall issue a new certificate of registration to the registrant, who shall maintain it with the old certificate of registration until the expiration date. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.61; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-28 Termination of registration; notice to board

Authority: IC 35-48-3-1 Affected: IC 35-48-3-6

Sec. 28. Termination of registration. The registration of any person shall terminate if and when such person dies, ceases legal existence, or discontinues business or professional practice. Any registrant who ceases legal existence or discontinues business or professional practice shall notify the Indiana Board of Pharmacy promptly of such fact. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.62; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-29 Transfer of registration

Authority: IC 35-48-3-1 Affected: IC 35-48-3-6

Sec. 29. Transfer of registration. No registration or any authority conferred thereby shall be assigned or otherwise transferred except upon such conditions as the Indiana Board of Pharmacy may specifically designate and then only pursuant to its written consent. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.63; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-30 Security requirements; approval of security system

Authority: IC 35-48-3-1 Affected: IC 35-48-3-7

Sec. 30. Security requirements generally. (a) All applicants and registrants shall provide and maintain effective controls and procedures to guard against theft and diversion of controlled substances. In order to determine whether a registrant has provided effective controls against diversion, the Indiana Board of Pharmacy shall use the security requirements set forth in Sections 3.72-3.76 *[856 IAC 2-3-31 – 856 IAC 2-3-35]* as standards for the physical security controls and operating procedures necessary to prevent diversion. Materials and construction which will provide a structural equivalent to the physical security controls set forth in Sections 3.72 *[856 IAC 2-3-31]*, 3.73 *[856 IAC 2-3-32]*, and 3.75 *[856 IAC 2-3-34]* may be used in lieu of the materials and construction described in those sections.

(b) Substantial compliance with the standards set forth in Sections 3.72-3.76 [856 IAC 2-3-31 – 856 IAC 2-3-35] may be deemed sufficient by the Indiana Board of Pharmacy after evaluation of the overall security system and needs of a registrant or applicant. In evaluating the overall security system of a registrant or applicant, the Indiana Board of Pharmacy may consider any of the following factors as it may deem relevant to the need for strict compliance with security requirements:

(1) The type of activity conducted (e.g., processing of bulk chemicals, preparing dosage forms, packaging, labeling, cooperative buying, etc.);

(2) The type and form of controlled substances handled (e.g., bulk liquids or dosage units usable powders or nonusable powders);

(3) The quantity of controlled substance handled;

(4) The location of the premises and the relationship such location bears on security needs;

(5) The type of building construction comprising the facility and the general characteristics of the building or buildings;

(6) The type of vault, safe, and secure enclosures or other storage system (e.g., automatic storage and retrieval system) used;

(7) The type of closures on vaults, safes, and secure enclosures;

(8) The adequacy of key control systems and/or combination lock control systems;

(9) The adequacy of electric detection and alarm systems, if any including use of supervised transmittal lines and standby power sources;

(10) The extent of unsupervised public access to the facility, including the presence and characteristics of perimeter fencing, if any;

(11) The adequacy of supervision over employees having access to manufacturing and storage areas;

(12) The procedures for handling business guests, visitors, maintenance personnel, and non-employee service personnel;

(13) The availability of local police protection or of the registrant's or applicant's security personnel, and;

(14) The adequacy of the registrant's or applicant's system for monitoring the receipt, manufacture, distribution, and disposition of controlled substances in its operations.

(c) When physical security controls become inadequate as a result of a controlled substance being transferred to a different schedule, or as a result of a non-controlled substance being listed on any schedule, or as a result of a significant increase in the quantity of controlled substances in the possession of the registrant during normal business operations, the physical security controls shall be expanded and extended accordingly. A registrant may adjust physical security controls within the requirements set forth in Sections 3.72-3.76 [856 IAC 2-3-31 – 856 IAC 2-3-35] when the need for such controls decreases as a result of a controlled substance being transferred to a different schedule, or a result of a controlled substance being removed from control, or as a result of a significant decrease in the quantity of controlled substances in the possession of the registrant during normal business operations.

(d) Any registrant or applicant desiring to determine whether a proposed security system substantially complies with, or is the structural equivalent of, the requirements set forth in Sections 3.72-3.76 [856 IAC 2-3-31 – 856 IAC 2-3-35] may submit any plans, blueprints, sketches or other materials regarding the proposed security system to the Indiana Board of Pharmacy.

(e) Approval by the Drug Enforcement Administration of any security system, proposed security system, plans, blueprints, sketches or other material as being in substantial compliance with the requirements as set forth in 301.72-301.76 of Title 21 of the Code of Federal Regulations shall be deemed in compliance with Sections 3.71 through 3.75 [856 IAC 2-3-30 – 856 IAC 2-3-34] of these regulations, where applicable.

(f) Physical security controls of locations registered under the Harrison Narcotic Act or the Narcotics Manufacturing Act of 1960 on April 20, 1971, shall be deemed to comply substantially with the standards set forth in Sections 3.71 [this section], 3.72 [856 IAC 2-3-31], 3.73 [856 IAC 2-3-32], and 3.75 [856 IAC 2-3-34]. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.71; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-31 Storage areas; security controls for nonpractitioners

Authority: IC 35-48-3-1 Affected: IC 35-48-3-7

Sec. 31. Physical security controls for nonpractitioners: Storage Areas. (a) Schedules I and II [856 IAC 2-2-2 and 856 IAC 2-2-3]. Raw materials, bulk materials awaiting further processing, and finished products which are controlled substances listed in Schedule I or II [856 IAC 2-2-2 or 856 IAC 2-2-3] shall be stored in one of the following secure storage areas:

(1) Where small quantities permit, a safe or steel cabinet.

(i) Which safe or steel cabinet shall have the following specifications or the equivalent: 30 man-minutes against surreptitious entry, 10 man-minutes against forced entry, 20 man-hours against lock manipulation, and 20 man-hours against radiological techniques.

(ii) Which safe or steel cabinet, if it weighs less than 750 pounds, is bolted or cemented to the floor or wall in such a way that it cannot be readily removed; and

(iii) Which safe or steel cabinet, if necessary, depending upon the quantities and type of controlled substances stored, is equipped with an alarm system which, upon attempted unauthorized entry, shall transmit a signal directly to a central protection company or a local or State police agency which has a legal duty to respond, or a 24-hour control station operated by the registrant, or such other protection as the Indiana Board of Pharmacy or the Drug Enforcement Administration may approve.

(2) A vault constructed before, or under construction on October 1, 1973, which is of substantial construction with a steel door, combination or key lock, and an alarm system; or

(3) A vault constructed after October 1, 1973:

(i) The walls, floors, and ceilings of which vault are constructed of at least 8 inches of reinforced concrete or other substantial masonry, reinforced vertically and horizontally with 1/2-inch steel rods tied 6 inches on center, or the structural equivalent to such reinforced walls, floors, and ceilings;

(ii) The door and frame unit of which vault shall conform to the following specifications or the equivalent; 30 manminutes against surreptitious entry, 10 man-minutes against forced entry, 20 man-hours against lock manipulation, and 20 man-hours against radiological techniques;

(iii) Which vault, if operations require it to remain open for frequent access, is equipped with "day-gate" which is selfclosing and self-locking, or the equivalent, for use during the hours of operation in which the vault door is open; (iv) The walls or perimeter of which vault are equipped with an alarm, which upon unauthorized entry shall transmit a signal directly to a central station protection company, or a local or State police agency which has a legal duty to respond or a 24-hour control station operated by the registrant or such other protection as the Indiana Board of Pharmacy or the Drug Enforcement Administration may approve, and, if necessary, holdup buttons at strategic points of entry to the perimeter area of the vault;

(v) The door of which vault is equipped with contact switches; and

(vi) Which vault has one of the following: complete electrical lacing of the walls, floor and ceilings; sensitive ultrasonic equipment within the vault; a sensitive sound accumulator system; or such other device designed to detect illegal entry as may be approved by the Indiana Board of Pharmacy or the Drug Enforcement Administration.

(b) Schedules III, IV, and V [856 IAC 2-2-4-856 IAC 2-2-6]. Raw materials, bulk materials awaiting further processing, and finished products which are controlled substances listed in Schedules III, IV, and V [856 IAC 2-2-4-856 IAC 2-2-6] shall be stored in one of the following secure storage areas:

(1) Where small quantities permit, a safe which complies with the requirements set forth in paragraph (a)(1) of this section;

(2) A vault which complies with the requirements set forth in either paragraph (a)(2) or (3) of this section or

(3) A building or area located within a building, which building or area:

(i) Has walls or perimeter fences of sufficient height and construction to provide security from burglary;

(ii) Has substantial doors which may be securely locked during non-working hours by a multiple-position combination or key lock;

(iii) Is equipped with an alarm which, upon unauthorized entry, shall transmit a signal directly to a central station protection company, or local or State police agency which has a legal duty to respond or a 24-hour control station operated by the registrant or such other protection as the Indiana Board of Pharmacy or the Drug Enforcement Administration may approve; and

(iv) In which all controlled substances are segregated from all other merchandise and kept under constant surveillance during normal business hours.

(c) Multiple storage areas. Where several types or classes of controlled substances are handled separately by the registrant or applicant for different purposes (e.g., returned goods, or goods in process), the controlled substances may be stored separately, provided that each storage area complies with the requirements set forth in this section.

(d) Accessibility to storage areas. The controlled substances storage areas shall be accessible only to an absolute minimum number of specifically authorized employees. When it is necessary for employee maintenance personnel, non-employee maintenance personnel, business guests, or visitors to be present in or pass through controlled substances storage areas, the registrant shall provide for adequate observation of the area by an employee specifically authorized in writing.

(e) Compliance with the requirements prescribed in Part 301, Section 301.72 of Title 21 of the Code of Federal Regulations, effective April 1, 1973, shall be deemed in compliance with this section. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.72; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-32 Manufacturing areas; security controls for nonpractitioners

Authority: IC 35-48-3-1 Affected: IC 35-48-3-7

Sec. 32. Physical security controls for nonpractitioners: Manufacturing areas. All manufacturing activities (including processing, packaging, and labeling) involving controlled substances listed in any schedule shall be conducted in accordance with the following:

(a) All in-process substances shall be returned to the controlled substances storage area at the termination of the process. If the process is not terminated at the end of a workday (except where a continuous process or other normal manufacturing operation should not be interrupted), the processing area or tanks, vessels, bins or bulk containers containing such substances shall be securely locked, with adequate security for the area or building. If such security requires an alarm, such alarm, upon unauthorized entry, shall transmit a signal directly to a central station protection company, or a local or State Police agency which has a legal duty to respond, or a 24-hour control station operated by registrant.

(b) Manufacturing activities with controlled substances shall be conducted in an area or areas of clearly defined limited access which is under surveillance by an employee or employees designated in writing as responsible for the area. "Limited access"

may be provided in the absence of physical dividers such as walls or partitions, by traffic control lines or restricted space designation. The employee designated as responsible for the area may be engaged in the particular manufacturing operation being conducted: Provided: that he is able to provide continuous surveillance of the area in order that unauthorized persons may not enter or leave the area without his knowledge.

(c) During the production of controlled substances, the manufacturing areas shall be accessible to only those employees required for efficient operation. When it is necessary for employee maintenance personnel, non-employee maintenance personnel, business guests, or visitors to be present in or pass through manufacturing areas during production of controlled substances, the registrant shall provide for adequate observation of the area by an employee specifically authorized in writing. (d) Compliance with the requirements prescribed in Part 301, Section 301.73 of Title 21 of the Code of Federal Regulations, effective April 1, 1973, shall be deemed in compliance with this section.

(Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.73; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-33 Additional security controls for nonpractitioners

Authority: IC 35-48-3-1 Affected: IC 35-48-3-7

Sec. 33. Other security controls for nonpractitioners: (a) Before distributing a controlled substance to any person who the registrant does not know to be registered to possess the controlled substance the registrant shall make a good faith inquiry either with the Indiana Board of Pharmacy or with the D.E.A. to determine that the person is registered to possess the controlled substance.

(b) The registrant shall design and operate a system to disclose to the registrant suspicious orders of controlled substances. The registrant shall inform the Indiana Board of Pharmacy of suspicious orders when discovered by the registrant. Suspicious orders include orders of unusual size, orders deviating substantially from a normal pattern, and orders of unusual frequency.

(c) The registrant shall notify in writing the Indiana Board of Pharmacy of any theft or significant loss of any controlled substances upon discovery of such theft or loss. Thefts must be reported whether or not the controlled substances are subsequently recovered and/or the responsible parties are identified and action taken against them.

(d) The registrant shall not distribute any controlled substance listed in Schedules II through V [856 IAC 2-2-3 - 856 IAC 2-2-6] as a complimentary sample to any potential or current customer (1) without the prior written request of the customer, (2) to be used only for satisfying the legitimate medical needs of patients of the customer, and (3) only in reasonable quantities. Such request must contain the name, address and state and federal registration number of the customer and the name and quantity of the specific controlled substance desired. The request shall be preserved by the registrant with other records of distribution of controlled substances. In addition, the requirements of Part 3.5 [856 IAC 2-3-24 - 856 IAC 2-3-26] hereof shall be complied with for any distribution of a controlled substance listed in Schedule II [856 IAC 2-2-3]. For purposes of this paragraph, the term "customer" includes a person to whom a complimentary sample of a substance is given in order to encourage the prescribing or recommending of the substance by the person.

(e) When shipping controlled substances, a registrant is responsible for selecting common or contract carriers which provide adequate security to guard against in-transit losses. When storing controlled substances in a public warehouse, a registrant is responsible for selecting a warehouseman which will provide adequate security to guard against storage losses; wherever possible, the registrant shall store controlled substances in a public warehouse which complies with the requirements set forth in Section 3.72 *[856 IAC 2-3-31].* In addition, the registrant shall employ precautions (e.g., assuring that shipping containers do not indicate that contents are controlled substances) to guard against storage or in-transit losses.

(f) When distributing controlled substances through agents (e.g., detailmen), a registrant is responsible for providing and requiring adequate security to guard against theft and diversion while the substances are being stored or handled by the agent or agents. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.74; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-34 Storage; security controls for practitioners

Authority: IC 35-48-3-1 Affected: IC 35-48-3-7

Sec. 34. Physical security controls for practitioners. (a) Controlled substances listed in Schedule I [856 IAC 2-2-2] shall be

stored in a securely locked substantially constructed cabinet.

(b) Controlled substances listed in Schedules II, III, IV, and V [856 IAC 2-2-3 – 856 IAC 2-2-6] shall be stored in a securely locked, substantially constructed cabinet. However, pharmacies and institutional practitioners (as defined) in Chapter I, 1.01 [856 IAC 2-1-1] may disperse such substances throughout the stock of noncontrolled substances in such a manner as to obstruct the theft or diversion of the controlled substances.

Controlled substances (institutional practitioners–ward and floor stock) listed in Schedule II [856 IAC 2-2-3] and narcotic drugs in Schedule III [856 IAC 2-2-4] shall be stored in a securely locked substantially constructed cabinet or device. Controlled substances (institutional practitioners ward and floor stock) listed in Schedules III, IV, and V [856 IAC 2-2-4–856 IAC 2-2-6] may be dispersed in their ward or floor stock in such a manner as to obstruct theft or diversion of these controlled substances.

(c) This section shall also apply to nonpractitioners authorized to conduct research or chemical analysis under another registration. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.75; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-3-35 Additional security controls for practitioners

Authority:	IC 35-48-3-1
Affected:	IC 35-48-3-7

Sec. 35. Other security controls for practitioners. (a) The registrant shall not employ as an agent or employee who has access to controlled substances any person who has had an application for registration denied, or has had his registration revoked, or has been convicted of a violation of State or Federal law relative to the manufacture, distribution, dispensing or possession of controlled substances.

(b) The registrant shall notify the Indiana Board of Pharmacy of the theft or significant loss of any controlled substances upon discovery of such loss or theft. (Indiana Board of Pharmacy; Reg 28, Ch III, Sec 3.76; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

Rule 4. Records and Inventories of Registrants

856 IAC 2-4-1 Records and inventories

Authority: IC 35-48-3-1 Affected: IC 35-48-3-7

Sec. 1. (a) Every registrant shall keep records and maintain inventories in conformance with the record keeping and inventorying requirements of federal law and regulation.

(b) For purposes of this section, "readily retrievable" means that certain records are kept by automatic data processing systems or other electronic or mechanized record keeping systems in such a manner that they can be separated out from all other records in a reasonable time and/or records are kept on which certain items are asterisked, red-lined, or in some manner visually identifiable apart from other items appearing on the records. Manufacturers, distributors, and research records or electronic data processing printouts shall be made available within five (5) working days after a request by the Indiana board of pharmacy for such records or information on controlled substances transactions.

(c) Each registered pharmacy shall maintain, for a period of two (2) years, its prescriptions of controlled substances by maintaining any of the following:

(1) Three (3) separate files as follows:

(A) A file for Schedule II drugs dispensed.

(B) A file for Schedules III, IV, and V drugs dispensed.

(C) A file for prescriptions for all other drugs dispensed.

(2) Two (2) separate files as follows:

(A) A file for all noncontrolled drugs dispensed.

(B) Another file for all controlled drugs dispensed in Schedules II, III, IV, and V. If this method is used, the prescriptions in the file for Schedules III, IV, and V must be stamped with the letter "C" in red ink, not less than one (1) inch high, in the lower right-hand corner.

(3) Two (2) separate files as follows:

(A) A file for Schedule II drugs dispensed.

(B) Another file for Schedules III, IV, and V drugs, including all other noncontrolled drugs dispensed. If this method is used, the prescriptions in the file of Schedules III, IV, and V drugs must be stamped with the letter "C" in red ink, not less than one (1) inch high, in the lower right-hand corner.

However, if a pharmacy employs an automated data processing system or other electronic record keeping system for prescriptions that permits identification by prescription number and retrieval of original documents by prescriber's name, patient's name, drug dispensed, and date filled, then the requirement to mark the hard copy prescription with a red "C" under subdivisions (2) and (3) is waived. (Indiana Board of Pharmacy; Reg 28, Ch IV, Sec 4.01; filed Jul 9, 1974, 9:29 a.m.: Unpublished; filed May 26, 2000, 8:52 a.m.: 23 IR 2504; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

Rule 5. Order Forms

856 IAC 2-5-1 Order form requirements

Authority:	IC 35-48-3-1
Affected:	IC 35-48-3-8

Sec. 1. Order Form Requirements Generally. Compliance with the requirements prescribed in section 308 of the Federal Controlled Substances Act (21 U.S.C. 828), and in Part 305 of Title 21 of the Code of Federal Regulations, effective April 1, 1973 shall be deemed compliance with the requirements of IC 1971, 35-24.1-3-7 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended. (Indiana Board of Pharmacy; Reg 28, Ch V, Sec 5.01; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

Rule 6. Issuance, Filling and Filing Prescriptions

856 IAC 2-6-1 Scope of rules governing prescriptions

Authority:	IC 35-48-3-1
Affected:	IC 35-48-3-9

Sec. 1. Scope of Part 6. Rules governing the issuance, filling and filing of prescriptions pursuant to IC 1971, 35-24.1-3-8 [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended, are set forth generally in that section and specifically by the sections of this part [856 IAC 2]. (Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.01; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-2 Persons entitled to issue prescriptions

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Authority: IC 35-48-3-1
Affected: IC 35-48-3-9
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Sec. 2. (a) A prescription for a controlled substance may be issued only by an individual practitioner who is:

(1) authorized to prescribe controlled substances by the state; and

(2) either registered or exempted from registration pursuant to 856 IAC 2-3-5(b) or 856 IAC 2-3-6.

(b) A prescription issued by an individual practitioner may be communicated to a pharmacist by an individual practitioner or a practitioner's authorized agent.

(c) Controlled substances prescriptions issued by individual practitioners in adjoining states to Indiana or other states are considered valid prescriptions if the practitioner issuing the prescription has a current and valid Drug Enforcement Administration certificate registration number. It is the pharmacist's responsibility as with all controlled substances prescriptions, to be sure beyond reasonable doubt in his or her professional judgment that the practitioner is issuing the prescription in good faith and has a valid Drug Enforcement Administration certificate of registration. (Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.02; filed Jul 9, 1974, 9:29 a.m.: Unpublished; readopted filed Nov 30, 2001, 11:00 a.m.: 25 IR 1343; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-

856070060RFA)

856 IAC 2-6-3 Purpose of prescription; prohibitions

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Authority: IC 35-48-3-1
Affected: IC 35-48-3-9
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Sec. 3. Purpose of issue of prescription. (a) A prescription for a controlled substance to be effective must be issued for a legitimate medical purpose in a reasonable quantity by an individual practitioner acting in the usual course of his professional practice. The responsibility for the proper prescribing and dispensing of controlled substances is upon the prescribing practitioner, but a corresponding responsibility rests with the pharmacist who fills the prescription. An order purporting to be a prescription issued not in the usual course of professional treatment or in legitimate and authorized research is not a prescription, within the meaning and intent of IC 1971, 35-24.1-3-8 *[Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.]* as amended, and the person knowingly filling such a purported prescription, as well as the person issuing it, shall be subject to the penalties provided for violations of the provisions of law relating to controlled substances.

(b) A prescription may not be issued in order for an individual practitioner to obtain controlled substances for supplying the individual practitioner for the purpose of general dispensing to patients.

(c) A prescription may not be issued for the dispensing of narcotic drugs listed in any schedule to a narcotic drug dependent person for the purpose of continuing his dependence upon such drugs in the course of conducting an authorized clinical investigation in the development of a narcotic addict rehabilitation program. (Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.03; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-4 Issuance of prescriptions; information required

Authority: IC 35-48-3-1 Affected: IC 35-48-3-9

Sec. 4. Manner of issuance of prescriptions. (a) All prescriptions for controlled substances shall be dated as of, and signed on, the day when issued and shall bear the full name and address of the patient, and the name, address, and Federal Controlled Substance registration number of the practitioner. A practitioner may sign a prescription in the same manner as he would sign a check or legal document (e.g., J.H. Smith or John H. Smith). Where an oral order is not permitted, prescriptions shall be written with ink or indelible pencil or typewriter and shall be manually signed by the practitioner. The prescriptions may be prepared by a secretary or agent for the signature of a practitioner, but the prescribing practitioner is responsible in case the prescription does not conform in all essential respects to the law and regulations. A corresponding liability rests upon the pharmacist who fills a prescription not prepared in the form prescribed by these regulations.

(b) An intern, resident, or foreign-trained physician exempted from registration under section 3.14(c) [856 IAC 2-3-5(c)], shall include on all prescriptions issued by him the Federal Controlled Substance registration number of the hospital or other institution and the special internal code number assigned to him by the hospital or other institution as provided in section 3.14(c) [856 IAC 2-3-5(c)], in lieu of the registration number of the practitioner required by this section. Each prescription shall have the name of the intern, resident, or foreign-trained physician stamped or printed on it, as well as the signature of the physician.

(c) An official exempted from registration under section 3.15 [856 IAC 2-3-6] shall include on all prescriptions issued by him, his branch of service or agency (e.g., "U.S. Army" or "Public Health Service") and his service identification number, in lieu of the Federal Controlled Substance registration number of the practitioner required by this section. The service identification number for a Public Health Service employee is his Social Security identification number. Each prescription shall have the name of the officer stamped, typed, or hand-printed on it, as well as the signature of the officer. (Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.04; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-5 Persons entitled to fill prescriptions

Authority: IC 35-48-3-1 Affected: IC 35-48-3-9

Sec. 5. Persons entitled to fill prescriptions. A prescription for controlled substances may only be filled by a pharmacist acting in the usual course of his professional practice and either registered individually or employed in a registered pharmacy or by a registered institutional practitioner. (*Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.05; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA*)

856 IAC 2-6-6 Dispensing of narcotics for maintenance purposes

Authority: IC 35-48-3-1 Affected: IC 35-48-3-9

Sec. 6. Dispensing of narcotic drugs for maintenance purposes. The administering or dispensing directly (but not prescribing) of narcotic drugs listed in any schedule to a narcotic drug dependent person for the purpose of treatment of his dependence upon such drugs in the course of conducting a clinical investigation authorized by State or Federal law in the development of a narcotic addict rehabilitation program shall be deemed to be within the meaning of the term "in the course of his professional practice or research" in IC 1971, 35-24.1-1-1(u) [Repealed by Acts 1976, P.L.148, SECTION 24; Acts 1977, P.L.26, SECTION 25. See IC 35-48.] as amended. (Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.06; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-7 Schedule II controlled substances; prescription required; exceptions

Authority: IC 35-48-3-1 Affected: IC 35-48

Sec. 7. (a) A pharmacist may dispense directly a controlled substance listed in Schedule II, which is a prescription drug as determined under the Federal Food, Drug, and Cosmetic Act, only pursuant to a written prescription signed by the prescribing individual practitioner, except as provided in subsection (d).

(b) An individual practitioner may administer or dispense directly a controlled substance listed in Schedule II in the course of his or her professional practice without a prescription subject to section 6 of this rule.

(c) An institutional practitioner may administer or dispense directly (but not prescribe) a controlled substance listed in Schedule II only pursuant to a written prescription signed by the prescribing individual practitioner or to an order for medication made by an individual practitioner that is dispensed for immediate administration to the ultimate user.

(d) In the case of an emergency situation, as defined by subsection (e), a pharmacist may dispense a controlled substance listed in Schedule II upon receiving oral authorization of a prescribing individual practitioner, provided the following:

(1) The quantity prescribed and dispensed is limited to the amount adequate to treat the patient during the emergency period (dispensing beyond the emergency period must be pursuant to a written prescription signed by the prescribing individual practitioner).

(2) The prescription shall be immediately reduced to writing by the pharmacist and shall contain all information required in section 4 of this rule, except for the signature of the prescribing individual practitioner.

(3) If the prescribing individual practitioner is not known to the pharmacist, he or she must make a reasonable effort to determine that the oral authorization came from a registered individual practitioner, which may include a callback to the prescribing individual practitioner using his or her phone number as listed in the telephone directory and/or other good faith efforts to assure his or her identity.

(4) Within seven (7) days after authorizing an emergency oral prescription, the prescribing individual practitioner shall cause a written prescription for the emergency quantity prescribed to be delivered to the dispensing pharmacist. In addition to conforming to the requirements of section 4 of this rule, the prescription shall have written on its face "Authorization for Emergency Dispensing", and the date of the oral order. The written prescription may be delivered to the pharmacist in person or by mail, but if delivered by mail, it must be postmarked within the seven (7) day period. Upon receipt, the dispensing pharmacist shall attach this prescription to the oral emergency prescription which had earlier been reduced to writing. The pharmacist shall notify the Indiana board of pharmacy if the prescribing individual fails to deliver a written prescription to him or her, failure of the pharmacist to do so shall void the authority conferred by this subdivision to dispense without a written prescription of a prescribing individual practitioner.

(e) For the purpose of authorizing an oral prescription of a controlled substance listed in Schedule II of IC 35-48 as amended, "emergency situation" means those situations in which the prescribing practitioner determines the following:

(1) That immediate administration of the controlled substance is necessary for proper treatment of the intended ultimate user.
 (2) That no appropriate alternative treatment is available, including administration of a drug that is not a controlled substance under Schedule II of IC 35-48 as amended.

(3) That it is not reasonably possible for the prescribing practitioner to provide a written prescription to be presented to the person dispensing the substance, prior to the dispensing.

(Indiana Board of Pharmacy; Reg 28, Ch VI,Sec 6.11; filed Jul 9, 1974, 9:29 a.m.: Unpublished; filed May 26, 2000, 8:52 a.m.: 23 IR 2505; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-8 Schedule II controlled substances; refilling prescriptions

Authority: IC 35-48-3-1 Affected: IC 35-48-3-9

Sec. 8. Refilling prescriptions–Schedule II [856 IAC 2-2-3]. The refilling of a prescription for a controlled substance listed in Schedule II [856 IAC 2-2-3] is prohibited. (Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.12; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-9 Schedule II controlled substances; partial filling of prescriptions

Authority: IC 35-48-3-1

Affected: IC 35-48-2-6; IC 35-48-3-9

Sec. 9. (a) The partial filling of a prescription for a controlled substance listed in Schedule II under IC 35-48-2-6, as amended, is permissible if the pharmacist is unable to supply the full quantity called for in a written or emergency oral prescription and makes a notation of the quantity supplied on the face of the written prescription (or written record of the emergency oral prescription). The remaining portion of the prescription may be filled within seventy-two (72) hours of the first partial filling; however, if the remaining portion is not or cannot be filled within the seventy-two (72) hour period, the pharmacist shall so notify the prescribing individual practitioner. No further quantity may be supplied beyond seventy-two (72) hours without a new prescription.

(b) A prescription for a Schedule II controlled substance written for patients in long term care facilities may be filled in partial quantities to include individual dosage units. For each partial filling, the dispensing pharmacist shall record on the back of the prescription (or on another appropriate record, uniformly maintained, and readily retrievable) the date of the partial filling, quantity dispensed, remaining quantity authorized to be dispensed, and the identification of the dispensing pharmacist. The total quantity of a Schedule II controlled substance dispensed in all partial fillings must not exceed the total quantity prescribed. A Schedule II prescription, for a patient in a long term care facility, shall be valid for a period not to exceed sixty (60) days from the issue date unless sooner terminated by the discontinuance of medication.

(c) A prescription for a Schedule II controlled substance written for a patient with a medical diagnosis documenting a terminal illness may be filled in partial quantities to include individual dosage units. If there is any question whether a patient may be classified as having a terminal illness, the pharmacist must contact the practitioner prior to partially filling the prescription. The pharmacist has a responsibility to assure that the controlled substance is for a terminally ill patient. The pharmacist must record on the prescription the patient is "terminally ill". A prescription that is partially filled and does not contain the notation "terminally ill" shall be deemed to have been filled in violation of this section. For each partial filling, the dispensing pharmacist shall record on the back of the prescription (or on another appropriate record, uniformly maintained and readily retrievable) the date of the partial filling, quantity dispensed, remaining quantity authorized to be dispensed, and the identification of the dispensing pharmacist. Prior to any subsequent partial filling, the pharmacist is to determine that the additional partial filling is necessary. The total quantity of a Schedule II controlled substance dispensed in all partial fillings must not exceed the total quantity prescribed. A Schedule II prescription for a patient with a medical diagnosis documenting a terminal illness shall be valid for a period not to exceed sixty (60) days from the issue date unless sooner terminated by the discontinuance of medication. *(Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.13; filed Jul 9, 1974, 9:29 a.m.: Unpublished; filed May 8, 1986, 9:55 a.m.: 9 IR 2205; filed Mar 31, 1992, 5:00 p.m.: 15 IR 1391; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)*

856 IAC 2-6-10 Schedule II controlled substances; label information; exceptions

Authority: IC 35-48-3-1 Affected: IC 35-48-3-9

Sec. 10. Labeling of substances. (a) The pharmacist filling a written or emergency oral prescription for a controlled substance listed in Schedule II *[856 IAC 2-2-3]* shall affix to the package a label showing date of filling, the pharmacy name and address, the serial number of the prescription, the name of the patient, the name of the prescribing practitioner, and directions for use and the cautionary statement, "Federal Law prohibits the transfer of this drug to any person other than the patient for whom it was prescribed", any others if any, contained in such prescription or required by law.

(b) The requirements of paragraph (a) of this section do not apply when a controlled substance listed in Schedule II [856 IAC 2-2-3] is prescribed for administration to an ultimate user who is institutionalized; Provided, That:

(1) Not more than 7-day supply of the controlled substance listed in Schedule II [856 IAC 2-2-3] is dispensed at one time;

(2) The controlled substance listed in schedule II [856 IAC 2-2-3] is not in the possession of the ultimate user prior to the administration; and

(3) The institution maintains appropriate safeguards and records regarding the proper administration, control, dispensing, and storage of the controlled substance listed in schedule II [856 IAC 2-2-3]; and

(4) The system employed by the pharmacist in filling a prescription is adequate to identify the supplier, the product, and the patient, and to set forth the directions for use and cautionary statements, if any, contained in the prescription or required by law.

(Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.14; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-11 Schedule II controlled substances; retention of prescriptions (Repealed)

Sec. 11. (Repealed by Indiana Board of Pharmacy; filed Nov 30, 2001, 11:00 a.m.: 25 IR 1344)

856 IAC 2-6-12 Schedules III and IV controlled substances

Authority:	IC 35-48-3-1
Affected:	IC 35-48-3-9

Sec. 12. (a) A pharmacist may dispense a controlled substance listed in Schedule III or IV under 856 IAC 2-2-4 or 856 IAC 2-2-5, which is a prescription drug as determined under the federal Food, Drug, and Cosmetic Act, only pursuant to either a written prescription signed by a prescribing individual practitioner or an oral prescription made by a prescribing individual practitioner or a practitioner's authorized agent and promptly reduced to writing by the pharmacist containing all information required in 856 IAC 2-6-4 *[section 4 of this rule]*, except for the signature of the prescribing individual practitioner.

(b) An individual practitioner may administer or dispense a controlled substance listed in Schedule III or IV under 856 IAC 2-2-4 or 856 IAC 2-2-5 in the course of his or her professional practice without a prescription, subject to 856 IAC 2-6-6 [section 6 of this rule].

(c) An institutional practitioner may administer or dispense directly (but not prescribe) a controlled substance listed in Schedule III or IV under 856 IAC 2-2-4 or 856 IAC 2-2-5 pursuant to a written prescription signed by a prescribing individual practitioner, or pursuant to an oral prescription made by a prescribing individual practitioner and promptly reduced to writing by the pharmacist (containing all information required in 856 IAC 2-6-4 *[section 4 of this rule]*, except for the signature of the prescribing individual practitioner), or pursuant to an order for medication made by an individual practitioner that is dispensed for immediate administration to the ultimate user, subject to 856 IAC 2-6-6 *[section 6 of this rule]*. (Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.21; filed Jul 9, 1974, 9:29 a.m.: Unpublished; readopted filed Nov 30, 2001, 11:00 a.m.: 25 IR 1343; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-13 Schedules III, IV, and V controlled substances; refilling prescriptions; retrievable information Authority: IC 35-48-3-1

Affected: IC 35-48-2; IC 35-48-3-9

Sec. 13. (a) No prescription for a controlled substance listed in Schedule III (IC 35-48-2-8), Schedule IV (IC 35-48-2-10), or Schedule V (IC 35-48-2-12) shall be filled or refilled more than six (6) months after the date on which such prescription was issued, and no such prescription shall be authorized to be refilled more than five (5) times.

(b) Each refill of a prescription shall be recorded by one (1) of the following methods:

(1) On the back of the original prescription and, if used, a uniformly maintained, readily retrievable record such as a medication record or patient profile.

(2) In the storage memory of an electronic data processing system if such board approved system is used in the pharmacy.

(c) The following prescription information shall be retrievable by using or entering the serial number of the prescription:

(1) The name (and strength if applicable) and dosage form of the controlled substance.

(2) The date on which the prescription was written or phoned and reduced to writing by the pharmacist.

(3) The date of original filling and the date or dates of all refills.

(4) A notation or notations for the original filling and each and every subsequent refilling sufficient to identify the dispensing pharmacist.

(5) The total number of refills originally authorized and remaining for each individual prescription.

If the pharmacist does nothing more than date and initial the prescription to indicate a refill has been dispensed, the pharmacist shall be deemed to have dispensed a refill for the full face amount (that is the originally prescribed amount) of the prescription.

(d) Additional refills for prescriptions for controlled substances listed in Schedule III (IC 35-48-2-8), Schedule IV (IC 35-48-2-10), or Schedule V (IC 35-48-2-12) may be added to the original prescription on an oral authorization transmitted to the pharmacist by the original prescribing practitioner providing the following conditions are met:

(1) The total quantity authorized does not exceed the original face amount of the prescription and five (5) total refills, and none of the refills is for more dose units or a larger quantity than the original face amount of the prescription.

(2) No dispensing takes place pursuant to the original prescription more than six (6) months after the date of the original issue of the prescription.

(3) The pharmacist receiving the oral authorization records that authorization on the reverse of the original prescription, or in a readily retrievable record, and the following information:

(A) The date of the authorization.

(B) The number of the dose units or quantity authorized.

(C) The number of additional refills authorized.

(D) The initials of the pharmacist receiving the oral authorization.

(e) The prescribing practitioner must execute a new and separate prescription for any additional quantities beyond the five (5) refill, six (6) month limitation. (Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.22; filed Jul 9, 1974, 9:29 a.m.: Unpublished; filed May 20, 1988, 9:30 a.m.: 11 IR 3564; filed Jul 5, 1995, 10:00 a.m.: 18 IR 2783; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-14 Schedules III, IV and V controlled substances; partial filling of prescriptions

Authority: IC 35-48-3-1

Affected: IC 35-48-3-9

Sec. 14. The partial filling of a prescription for a controlled substance listed in Schedule III, IV, or V in the Controlled Substance Act, IC 35-48 as amended is permissible, provided that:

(a) each partial filling is recorded in the same manner as a refilling,

(b) the total quantity dispensed pursuant to an individual prescription including the original and all subsequent partial refills does not exceed the total quantity prescribed, and

(c) no dispensing occurs more than six (6) months after the date on which the prescription was issued.

(Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.23; filed Jul 9, 1974, 9:29 am: Unpublished; filed May 20, 1988, 9:30 am: 11 IR 3565; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-15 Schedules III and IV controlled substances; label information; exceptions

Authority: IC 35-48-3-1 Affected: IC 35-48-3-9 Sec. 15. Labeling of substances. (a) The pharmacist filling a prescription for a controlled substance listed in Schedule III or IV [856 IAC 2-2-4 or 856 IAC 2-2-5] shall affix to the package a label showing the pharmacy name and address, the serial number and date of initial filling, the name of the patient, the name of the practitioner issuing the prescription, and directions for use and cautionary statement, "Federal Law prohibits the transfer of this drug to any person other than the patient for whom it was prescribed", and other if any, contained in such prescription as required by law.

(b) The requirements of paragraph (a) of this section do not apply when a controlled substance listed in Schedule III or IV [856 IAC 2-2-4 or 856 IAC 2-2-5] is prescribed for administration to an ultimate user who is institutionalized: Provided, That:

(1) Not more than a 34-day supply or 100 dosage units, whichever is less, of the controlled substance listed in Schedule III or IV [856 IAC 2-2-4 or 856 IAC 2-2-5] is dispensed at one time;

(2) The controlled substance listed in schedule III or IV [856 IAC 2-2-4 or 856 IAC 2-2-5] is not in the possession of the ultimate user prior to administration;

(3) The institution maintains appropriate safeguards and records, the proper administration, control, dispensing, and storage of the controlled substance listed in Schedule III or IV [856 IAC 2-2-4 or 856 IAC 2-2-5]; and

(4) The system employed by the pharmacist in filling a prescription is adequate to identify the supplier, the product and the patient, and to set forth the directions for use and cautionary statements, if any, contained in the prescription or required by law.

(Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.24; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-16 Schedules III and IV controlled substances; retention of prescriptions

Authority: IC 35-48-3-1 Affected: IC 35-48-3-9

Sec. 16. Filing prescriptions. All prescriptions for controlled substances listed in Schedules III and IV [856 IAC 2-2-4 or 856 IAC 2-2-5] shall be kept in accordance with section 4.01 [856 IAC 2-4-1] of these regulations. (Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.25; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-17 Schedule V controlled substances; prescription requirements; refilling; exceptions

Authority: IC 35-48-3-1 Affected: IC 35-48-3-9

Sec. 17. Requirement of prescription. (a) A pharmacist may dispense a controlled substance listed in Schedule V [856 IAC 2-2-6] pursuant to a prescription as required for controlled substances listed in Schedules III and IV [856 IAC 2-2-4 and 856 IAC 2-2-5] in section 6.21 [856 IAC 2-6-12]. A prescription for a controlled substance listed in Schedule V [856 IAC 2-2-6] may be refilled only as expressly authorized by the prescribing individual practitioner on the prescription; if no such authorization is given, the prescription may not be refilled. A pharmacist dispensing such substance pursuant to a prescription shall label the substance in accordance with section 6.24 [856 IAC 2-6-15] and file the prescription in accordance with section 6.25 [856 IAC 2-6-16].

(b) An individual practitioner may administer or dispense a controlled substance listed in Schedule V [856 IAC 2-2-6] in the course of his professional practice without a prescription, subject to section 6.24 [856 IAC 2-6-15].

(c) An institutional practitioner may administer or dispense directly (but not prescribe) a controlled substance listed in Schedule V [856 IAC 2-2-6] only pursuant to a written prescription signed by the prescribing individual practitioner, or pursuant to an oral prescription made by a prescribing individual practitioner and promptly reduced to writing by the pharmacist (containing all information required in section 6.04 [856 IAC 2-6-4] except for the signature of the prescribing individual practitioner), or pursuant to an order for medication made by an individual practitioner which is dispensed for immediate administration to the ultimate user, subject to section 6.24 [856 IAC 2-6-15]. (Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.31; filed Jul 9, 1974, 9:29 am: Unpublished; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA)

856 IAC 2-6-18 Dispensing without prescription; delivery of devices

Authority: IC 35-48-3-1 Affected: IC 35-48-3-9

Sec. 18. (a) A controlled substance listed in Schedule V in the Controlled Substance Act, IC 35-48 which does not require a prescription under federal, state or local law or a device known as a hypodermic syringe and/or needle for human use may be dispensed by a pharmacist without a prescription to a purchaser at retail, provided that:

(1) such dispensing is made only by a pharmacist and not by a non-pharmacist employee even if under the direct supervision of a pharmacist (although after the pharmacist has fulfilled his professional and legal responsibilities set forth in his *[sic.]* section, the actual cash, credit transaction, or delivery may be completed by a non-pharmacist);
(2) no more than:

(i) 240 cc. (8 ounces) or 48 dosage units of any substance containing opium;

(ii) 120 cc. (4 ounces) or 24 dosage units of any other substance nor more than 48 dosage units may be dispensed at retail to the same purchaser in any given 48-hour period;

(3) the purchaser is at least eighteen (18) years of age. However, if the item being purchased is a device known as a hypodermic syringe and/or needle for human use, the age restriction shall not apply;

(4) the pharmacist requires every purchaser of a controlled substance or device as described in 856 IAC 2-6-18(a) not known to the pharmacist to furnish suitable identification (including proof of age where appropriate); and

(5) separate bound record books for dispensing of:

(i) controlled substances; and

(ii) devices under this section;

are maintained by the pharmacist. These books shall contain the name and address of the purchaser, the name and quantity of controlled substance or devices purchased, the date of each purchase, and the name or initials of the pharmacist who dispensed the substance or devices to the purchaser these books shall be maintained in accordance with the recordkeeping requirements of 856 IAC 2-4-1.

(b) Delivery of devices, as described above, to inpatients of institutions is exempt from this section.

(c) The delivery of a device known as a hypodermic syringe-needle other than by a pharmacist in a licensed pharmacy or a licensed practitioner in his lawful place of practice is prohibited. (*Indiana Board of Pharmacy; Reg 28, Ch VI, Sec 6.32; filed Jul 9, 1974, 9:29 am: Unpublished; filed May 20, 1988, 9:30 am: 11 IR 3565; readopted filed Nov 14, 2001, 2:45 p.m.: 25 IR 1341; readopted filed Oct 4, 2007, 3:33 p.m.: 20071031-IR-856070060RFA*)

Rule 7. Limited Permits

856 IAC 2-7-1 Application

Authority: IC 35-48-3-2 Affected: IC 35-48-3-2

Sec. 1. (a) A humane society, animal control agency, or governmental entity that intends to operate an animal shelter or other animal impounding facility for the purpose of buying, possessing, and using drugs authorized by IC 35-48-3-2 shall apply for a limited permit in the form and manner required by the board.

(b) The applicant shall provide the following:

(1) Name and address of the facility.

(2) Type of facility.

(3) Documentation describing the ownership of the facility.

(4) Fees set by the board in this rule.

(5) Information about the substances that the facility intends to administer.

(6) Written policies relating to storage, security, and procedures for access, handling, and administration of drugs.

(7) Proof that the employees of the applicant who will handle a controlled substance are sufficiently trained to use and administer the controlled substance.

(8) Proof that a licensed Indiana veterinarian holding a valid Indiana controlled substances registration and federal DEA registration has been retained to provide technical advice to the facility.

(c) No humane society, animal control agency, or governmental entity that intends to operate an animal shelter or other animal impounding facility for the purpose of buying, possessing, and using drugs authorized by IC 35-48-3-2 shall engage in any activity for which a permit is required until the permit is granted by the board. *(Indiana Board of Pharmacy; 856 IAC 2-7-1; filed Aug 21, 2003, 4:45 p.m.: 27 IR 181)*

856 IAC 2-7-2 Permit fees

Authority:	IC 35-48-3-2
Affected:	IC 35-48-3-2

Sec. 2. The board shall charge and collect the following fees:

(1) Application for a limited permit, fifty dollars (\$50).

(2) Annual renewal of limited permit, twenty-five dollars (\$25).

(Indiana Board of Pharmacy; 856 IAC 2-7-2; filed Aug 21, 2003, 4:45 p.m.: 27 IR 182)

856 IAC 2-7-3 Renewal of permit

Authority: IC 35-48-3-2 Affected: IC 35-48-3-2

Sec. 3. The renewal of the limited permits issued under this section shall be on the same schedule as other humane societies, animal control agencies, or governmental entities that hold controlled substance registrations issued by the board. (Indiana Board of Pharmacy; 856 IAC 2-7-3; filed Aug 21, 2003, 4:45 p.m.: 27 IR 182)

856 IAC 2-7-4 Storage, handling, and use of controlled substances

Authority: IC 35-48-3-2 Affected: IC 35-48-3-2

Sec. 4. (a) Limited permit holders and their agents, representatives, and employees must comply with the requirements of this rule for the storage and handling of controlled substances.

(b) All facilities at which controlled substances are stored, handled, or used shall:

(1) be of suitable size and construction to facilitate cleaning, maintenance, and proper operations;

(2) have storage areas large enough to provide adequate lighting, ventilation, temperature, sanitation, humidity, space, equipment, and security conditions;

(3) have a quarantine area for storage of controlled substances that are:

(A) outdated, damaged, deteriorated, misbranded, or adulterated; or

(B) in immediate or sealed secondary containers that have been opened;

(4) be maintained in a clean and orderly condition; and

(5) be free from infestation by insects, rodents, birds, or vermin of any kind.

(c) All facilities used for storage of controlled substances by registrants under this section shall comply with the security requirements as provided by 856 IAC 2-3-31.

(d) All controlled substances shall be stored at appropriate temperatures and under appropriate conditions in accordance with requirements, if any, in the labeling of such controlled substances or with requirements in the current edition of an official compendium of drug information.

(e) If no storage requirements are established for a controlled substance, the controlled substance may be held at a controlled room temperature, as defined in an official compendium, to help ensure that its identity, strength, quality, and purity are not adversely affected.

(f) Controlled substances that are outdated, damaged, deteriorated, misbranded, or adulterated shall be quarantined and physically separated from other controlled substances until they are destroyed by a designated agent of the board or returned to their supplier.

(g) Any controlled substance whose immediate or sealed outer or sealed secondary containers have been opened or used shall be:

(1) identified as such; and

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(2) quarantined and physically separated from other controlled substances until they are either destroyed by a designated agent of the board or returned to the supplier.

(h) Limited permit holders shall establish and maintain inventories and records of all controlled substances stored or used at the facility.

(i) Inventories and records shall be made available for inspection and photocopying by any authorized official of any governmental agency charged with enforcement of this rule for a period of two (2) years following disposition of the controlled substances.

(j) Records described in this section that are kept at the inspection site or that can be immediately retrieved by computer or other electronic means shall be readily available for authorized inspection during the retention period. Records kept at a central location apart from the inspection site and not electronically retrievable shall be made available for inspection within two (2) working days of a request by an authorized official of any governmental agency charged with enforcement of this rule. (Indiana Board of Pharmacy; 856 IAC 2-7-4; filed Aug 21, 2003, 4:45 p.m.: 27 IR 182)

856 IAC 2-7-5 Training of staff

Authority:	IC 35-48-3-2
Affected:	IC 35-48-3-2

Sec. 5. (a) Only employees of the limited permit holder are eligible for training to store, handle, and use controlled substances. Volunteers are prohibited from storing, handling, or using controlled substances.

(b) The following training is required:

(1) Completion of a comprehensive training program approved by the controlled substance advisory committee.

(2) Any additional training as required by the supervising veterinarian or site administrator.

(c) A veterinarian licensed to practice in Indiana, holding a valid Indiana controlled substances registration and federal DEA registration, must verify in writing that the employee has been trained adequately to store, handle, or use controlled substances. The written verification must be maintained at the facility in a reasonably retrievable manner.

(d) The limited permit holder or site administrator shall maintain documentary proof of training in a reasonably retrievable manner at the facility for review by an authorized official of any governmental agency charged with enforcement of this rule. (Indiana Board of Pharmacy; 856 IAC 2-7-5; filed Aug 21, 2003, 4:45 p.m.: 27 IR 183)

856 IAC 2-7-6 Protocol for administration of controlled substances

Authority: IC 35-48-3-2 Affected: IC 35-48-3-2

Sec. 6. In the event the consulting veterinarian is not physically present during the administration of controlled substances by employees of the limited permit holder, the veterinarian shall be available for consultation by telephonic or other electronic device. (Indiana Board of Pharmacy; 856 IAC 2-7-6; filed Aug 21, 2003, 4:45 p.m.: 27 IR 183)

856 IAC 2-7-7 Limitations on permit

Authority: IC 35-48-3-2 Affected: IC 35-48-3-2

Sec. 7. (a) Except as provided in subsection (b), only controlled substances for which the humane society, animal control agency, or governmental entity has received a permit may be stored, handled, and used at the facility.

(b) A licensed veterinarian who stores, handles, or uses controlled substances at the humane society, animal control agency, or governmental entity other than those authorized under the facility's limited permit, must apply for and obtain a controlled substance registration for the facility in the veterinarian's name.

(c) The veterinarian who holds the registration noted in subsection (b) is responsible for the proper storage, handling, and use of the controlled substances authorized for use under the veterinarian's controlled substance registration. (Indiana Board of Pharmacy; 856 IAC 2-7-7; filed Aug 21, 2003, 4:45 p.m.: 27 IR 183)

*