

Practitioner's Manual

An Informational Outline of the Controlled Substances Act

2006 Edition

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This manual has been prepared by the Drug Enforcement Administration, Office of Diversion Control, to assist practitioners (physicians, dentists, veterinarians, and other registrants authorized to prescribe, dispense, and administer controlled substances) in their understanding of the Federal Controlled Substances Act and its implementing regulations as they pertain to the practitioner's profession.

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Opioid (Narcotic) Addiction Treatment Programs
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SECTION I - INTRODUCTION

This practitioner's manual is intended to summarize and explain the basic requirements for prescribing, administering, and dispensing controlled substances under the Controlled Substances Act (CSA), 21 USC 801-890, and the DEA regulations, Title 21, Code of Federal Regulations (CFR), Parts 1300 to 1316. Pertinent citations to the law and regulations are included in this manual.

Printed copies of the CFR and the complete regulations implementing the CSA may be obtained from:

Superintendent of Documents U.S. Government Printing Office Washington, D.C. 20402

Both the CFR and the *Federal Register* (which includes proposed and final regulations implementing the CSA) are available on the Internet through the U.S. Government Printing Office (GPO) website. This website, which provides information by section, citation and keywords, can be accessed at:

www.gpoaccess.gov/cfr/index.html

Unofficial copies of pertinent CFR citations may be found at:

www.DEAdiversion.usdoj.gov

This practitioner's manual may also be found on the Internet at DEA's Web Site (under "publications"):

www.DEAdiversion.usdoj.gov

Should any pertinent provisions of the law or regulations be modified in the future, DEA will issue a revised electronic version of this document, which will be published on the DEA Diversion Website.

If you encounter errors in this document, please notify:

Editor, DEA Practitioner's Manual c/o DEA, Office of Diversion Control Liaison and Policy Section Washington, D.C. 20537

Inquiries regarding topics within this document may be addressed to your local DEA field office (listed in Appendix E) or the address above.

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This Document is Authorized for Public Dissemination

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Message from the Administrator

The Drug Enforcement Administration is pleased to provide this updated edition of the 1990 Practitioner's Manual to assist you in understanding your responsibilities under the Controlled Substances Act (CSA) and its implementing regulations. This manual will help answer questions that you may encounter in your practice and provide guidance in complying with federal requirements.

DEA remains committed to the 2001 Balanced Policy of promoting pain relief and preventing abuse of pain medications. In enforcing the CSA, it is DEA's responsibility to ensure drugs are not diverted for illicit purposes. Unfortunately, this country is now experiencing an alarming prescription drug abuse problem:

- Today, more than 6 million Americans are abusing prescription drugs—that is more than the number of Americans abusing cocaine, heroin, hallucinogens, and inhalants, combined.
- Researchers from the Centers for Disease Control and Prevention report that opioid prescription painkillers now cause more drug overdose deaths than cocaine and heroin combined.
- Today more new drug users have begun abusing pain relievers (2.4 million) than marijuana (2.1 million) or cocaine (1.0 million).

It is more important now than ever to be vigilant in preventing the diversion and abuse of controlled substances. This manual will help you do that by listing some safeguards you can take to prevent such diversion. It also explains registration, recordkeeping, and valid prescription requirements.

As a practitioner, your role in the proper prescribing, administering, and dispensing of controlled substances is critical to patients' health and to safeguarding society against the diversion of controlled substances. DEA is committed to working jointly with the medical community to ensure that those in need are cared for and that legitimate controlled substances are not being diverted for illegal use.

Karen P. Tandy Administrator September 2006

Preface

The Drug Enforcement Administration (DEA) was established in 1973 to serve as the primary federal agency responsible for the enforcement of the Controlled Substances Act (CSA). The CSA sets forth the federal law regarding both illicit and licit (pharmaceutical) controlled substances. With respect to pharmaceutical controlled substances, DEA's statutory responsibility is twofold: to prevent diversion and abuse of these drugs while ensuring an adequate and uninterrupted supply is available to meet the country's legitimate medical, scientific, and research needs. In carrying out this mission, DEA works in close cooperation with state and local authorities and other federal agencies.

Under the framework of the CSA, the DEA is responsible for ensuring that all controlled substance transactions take place within the "closed system" of distribution established by Congress. Under this "closed system," all legitimate handlers of controlled substances – manufacturers, distributors, physicians, pharmacies, and researchers – must be registered with DEA and maintain strict accounting for all distributions.

To carry out DEA's mission effectively, this 2006 Practitioner's Manual seeks to aid DEA registrants in complying with the CSA and its implementing regulations. The DEA understands that it can best serve the public interest by working with practitioners to prevent diversion of legal pharmaceutical controlled substances into the illicit market.

The federal controlled substances laws are designed to work in tandem with state controlled substance laws. Toward this same goal, DEA works in close cooperation with state professional licensing boards and state and local law enforcement officials to ensure that pharmaceutical controlled substances are prescribed, administered, and dispensed for legitimate medical purposes in accordance with federal and state laws. Within this cooperative framework, the majority of investigations into possible violations of the controlled substances laws are carried out by state authorities. However, DEA also conducts investigations into possible violations of federal law as circumstances warrant.

In the event a state board revokes the license of a practitioner, the DEA will take action and request a voluntary surrender of the practitioner's DEA registration. If the practitioner refuses to voluntarily surrender the registration, the DEA will pursue administrative action to revoke the DEA registration. The DEA may also pursue judicial action if there is sufficient evidence of illegal distribution or significant recordkeeping violations. All such actions are intended to deny the practitioner the means to continue to divert or abuse controlled substances as well as to protect the health and safety of the public and the practitioner.

The DEA is authorized under federal law to pursue legal action in order to prevent the diversion of controlled substances and protect the public safety. A lack of compliance may result in a need for corrective action, such as administrative action (that is, Letter of Admonition, an informal hearing or "order to show cause"), or in extreme cases, civil, or criminal action.

SECTION II – GENERAL REQUIREMENTS

Schedules of Controlled Substances

The drugs and other substances that are considered controlled substances under the CSA are divided into five schedules. A complete list of the schedules is published annually on an updated basis in the DEA regulations, Title 21 of the Code of Federal Regulations, Sections 1308.11 through 1308.15. Substances are placed in their respective schedules based on whether they have a currently accepted medical use in treatment in the United States and their relative abuse potential and likelihood of causing dependence when abused. Some examples of the drugs in each schedule are outlined below.

IMPORTANT NOTE:

All drugs listed in Schedule I have no currently accepted medical use in treatment in the United States and therefore may not be prescribed, administered, or dispensed for medical use. In contrast, drugs listed in Schedules II through V all have some accepted medical use and therefore may be prescribed, administered, or dispensed for medical use.

Schedule I Substances

Substances in this schedule have no currently accepted medical use in treatment in the United States, a lack of accepted safety for use under medical supervision, and a high potential for abuse.

Some examples of substances listed in Schedule I are: heroin; lysergic acid diethylamide (LSD); marijuana (cannabis); peyote; methaqualone; and methylene-dimethoxy-methamphetamine ("ecstasy").

The CSA allows for bona fide research with controlled substances in Schedule I, provided that the FDA has determined the researcher to be qualified and competent, and provided further that the FDA has determined the research protocol to be meritorious. Researchers who meet these criteria must obtain a separate registration to conduct research with a Schedule I controlled substance.

Schedule II Substances

Substances in this schedule have a high potential for abuse with severe psychological or physical dependence.

Examples of single entity Schedule II narcotics include morphine, codeine, and opium. Other Schedule II narcotic substances and their common name brand products include: hydromorphone (Dilaudid®), methadone (Dolophine®), meperidine (Demerol®), oxycodone (OxyContin®), and fentanyl (Sublimaze® or Duragesic®). Examples of Schedule II stimulants include amphetamine (Dexedrine® or Adderall®), methamphetamine (Desoxyn®), and methylphenidate (Ritalin®). Other Schedule II substances include: cocaine, amobarbital, glutethimide, and pentobarbital.

Schedule III Substances

Substances in this schedule have a potential for abuse less than substances in Schedules I or II.

Examples of Schedule III narcotics include combination products containing less than 15 milligrams of hydrocodone per dosage unit (i.e., Vicodin®) and products containing not more than 90 milligrams of codeine per dosage unit (i.e., Tylenol with codeine®).

Examples of Schedule III non-narcotics include benzphetamine (Didrex®), phendimetrazine, dronabinol (Marinol®), ketamine, and anabolic steroids such as oxandrolone (Oxandrin®).

Schedule IV Substances

Substances in this schedule have a lower potential for abuse relative to substances in Schedule III.

Examples of a Schedule IV narcotics include proposyphene (Darvon® and Darvocet-N 100®).

Other Schedule IV substances include alprazolam (Xanax®), clonazepam (Klonopin®), clorazepate (Tranxene®), diazepam (Valium®), lorazepam (Ativan®), midazolam (Versed®), temazepam (Restoril®), and triazolam (Halcion®).

Schedule V Substances

Substances in this schedule have a lower potential for abuse relative to substances listed in Schedule IV and consist primarily of preparations containing limited quantities of certain narcotic and stimulant drugs. These are generally used for antitussive, antidiarrheal and analgesic purposes.

Examples include cough preparations containing not more than 200 milligrams of codeine per 100 milliliters or per 100 grams (Robitussin AC®, and Phenergan with Codeine®).

Registration Requirements

Under the CSA, the term "practitioner" is defined as a physician, dentist, veterinarian, scientific investigator, pharmacy, hospital, or other person licensed, registered, or otherwise permitted, by the United States or the jurisdiction in which the practitioner practices or performs research, to distribute, dispense, conduct research with respect to, administer, or use in teaching or chemical analysis, a controlled substance in the course of professional practice or research. Every person or entity that handles controlled substances <u>must</u> be registered with DEA or be exempt by regulation from registration.

The DEA registration grants practitioners federal authority to handle controlled substances. However, the DEA registered practitioner may only engage in those activities that are authorized under state law for the jurisdiction in which the practice is located. When federal law or regulations differ from state law or regulations, the practitioner is required to abide by the more stringent aspects of both the federal and state requirements. In many cases, state law is more stringent than federal law, and must be complied with in addition to federal law. Practitioners should be certain they understand their state as well as DEA controlled substance regulations.

Application for Registration

To obtain a DEA registration, a practitioner must apply using a DEA Form 224. Applicants may submit the form by hard copy or on-line. Complete instructions accompany the form. To obtain the application, DEA may be contacted at:

- www.DEAdiversion.usdoj.gov (DEA Diversion Internet Web Site)
- any DEA field office (see listing in Appendix E of this manual)
- DEA Headquarters' Registration Section in Washington, D.C. at 1-800-882-9539 (Registration Call Center)

The DEA Form-224 may be completed on-line or in hard copy and mailed to:

Drug Enforcement Administration Registration Unit Central Station P.O. Box 28083 Washington, D.C. 20038-8083

A sample DEA Form 224 – New Application for Registration, is located at Appendix H, DEA Forms.

Certificate of Registration

The DEA Certificate of Registration (DEA Form 223) must be maintained at the registered location in a readily retrievable manner and kept available for official inspection.

The CSA requires that a separate registration be obtained for each principal place of business or professional practice where controlled substances are manufactured, distributed, or dispensed. DEA has historically provided an exception that a practitioner who is registered at one location, but also practices at other locations, is not required to register separately for any other location at which controlled substances are only prescribed. If the practitioner maintains supplies of controlled substances, administers, or directly dispenses controlled substances at the separate location the practitioner must obtain a separate DEA registration for that location. The exception applies only to a secondary location within the same state in which the practitioner maintains his/her registration. DEA individual practitioner registrations are based on state authority to dispense or conduct research with respect to controlled substances. Since a DEA registration is based on a state license, it cannot authorize controlled substance dispensing outside that state. Hence, the separate registration exception applies only to locations within the same state in which practitioner shave their DEA registrations.

A duplicate Certificate of Registration may be requested on-line. It appears on DEA's website, www.DEAdiversion.usdoj.gov, as follows:

	Internet of Justice Enforcement Administration DIVERSION CONTROL PROGRAM gistration Certificate Duplicate
	DEA Form 223 Duplicate Certificate Login:
DEA Number (R	equired - Not Case Sensitive)
If "Smith, John Q I If "Smith's, Pharm	your registration. Example: MD" is on your registration, then enter: Smith acy" is on your registration, then enter: Smith's acy" (no comma) is on your registration, n's Pharmacy
SSN (Required	if given on application)
Tax ID (Required	if given on application)
	ved your registration recently, your duplicate certificate may not xpire date, as some processing time is required.
	Login

Registration Renewals

Practitioner registrations must be renewed every three years. Renewal registrations use DEA Form 224a, Renewal Application for DEA Registration (see example at Appendix H, DEA Forms). The cost of the registration is indicated on the application form.

A renewal application is sent to the registrant approximately 45 days before the registration expiration date. The renewal application is sent to the address listed on the current registration certificate. If the renewal form is not received within 30 days before the expiration date of the current registration, the practitioner should contact the DEA registration office for their state, or DEA Headquarters at 1-800-882-9539, and request a renewal registration form.

The registration renewal application may be completed on-line at www.DEAdiversion.usdoj.gov, or in hard copy and mailed to:

Drug Enforcement Administration Registration Unit Central Station P.O. Box 28083 Washington, D.C. 20038-8083



Drug Registration > ODWIF

Registration Applications

Office of Diversion Control Web Interactive Forms (ODWIF)

RENEWAL APPLICATIONS

<u>Log-in to Begin</u> <u>Renewal Process</u>	Retail Pharmacy, Hospital/Clinic, Practitioner, Teaching Institution, or Mid-Level Practitioner, Manufacturer, Distributor, Researcher, Analytical Laboratory, Importer, Exporter, Domestic Chemicals
<u>Obtain Receipt</u>	This link may be used ONLY if you have previously submitted a Renewal Application through this tool and need an additional receipt.
<u>Duplicate</u> <u>Certificate</u>	On-line tool to request certificates for additional, misplaced, illegible, or destroyed originals.

MINIMUM ON-LINE REQUIREMENTS

The DEA Forms listed below are for those applying to DEA for a controlled substance registration. Data will be entered through a **secure connection** to the **ODWIF** on-line web application system. **Your web browser must support 128-bit encryption.**

You will need to have the following information handy in order to complete the form:

- Tax ID number and/or Social Security Number
- State Controlled Substance Registration Information
- State Medical License Information
- Credit Card (VISA, MasterCard, Discover or American Express)

The ODWIF system can only process credit card transactions at this time. If you are paying by check, you will need to <u>use the PDF version of the form</u>, then print and mail the form to the address listed on the form.

Change of Business Address

A practitioner who moves to a new physical location must request a modification of registration. A modification of registration can be requested on-line at www.DEAdiversion.usdoj.gov or in writing to the DEA field office responsible for that state. If the change in address involves a change in state, the proper state issued license and controlled substances registration must be obtained prior to the approval of modification of the federal registration. If the modification is approved, DEA will issue a new certificate of registration and, if requested, new Schedule II order forms (DEA Form-222, Official Order Form). A Renewal Application for Registration (DEA Form-224a) will only be sent to the registered address on file with DEA. It will not be forwarded.

Termination of Registration

Any practitioner desiring to discontinue business activities with respect to controlled substances must notify the nearest DEA field office (see Appendix E) in writing. Along with the notification of termination of registration, the practitioner should send the DEA Certificate of Registration and any unused Official Order Forms (DEA Form-222) to the nearest DEA field office.

Denial, Suspension or Revocation of Registration

Under the CSA, DEA has the authority to deny, suspend, or revoke a DEA registration upon a finding that the registrant has:

- 1. Materially falsified any application filed
- 2. Been convicted of a felony relating to a controlled substance or a List I chemical
- 3. Had their state license or registration suspended, revoked, or denied
- 4. Committed an act which would render the DEA registration inconsistent with the public interest
- 5. Been excluded from participation in a Medicaid or Medicare program

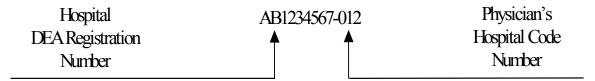
In determining the public interest, the CSA states the following factors are to be considered:

- 1. The recommendation of the appropriate state licensing board or professional disciplinary authority
- 2. The applicant's experience in dispensing or conducting research with respect to controlled substances
- 3. The applicant's conviction record under federal or state laws relating to the manufacture, distribution, or dispensing of controlled substances
- 4. Compliance with applicable state, federal, or local laws relating to controlled substances
- 5. Such other conduct which may threaten the public health and safety

Practitioner's Use of a Hospital's DEA Registration Number

Practitioners (e.g., intern, resident, staff physician, mid-level practitioner) who are agents or employees of a hospital or other institution may, when acting in the usual course of business or employment, administer, dispense, or prescribe controlled substances under the registration of the hospital or other institution in which they are employed, provided that:

- 1. The dispensing, administering, or prescribing is in the usual course of professional practice
- 2. Practitioners are authorized to do so by the state in which they practice
- 3. The hospital or institution has verified that the practitioner is permitted to dispense, administer or prescribe controlled substances within the state
- 4. The practitioner acts only within the scope of employment in the hospital or institution
- 5. The hospital or institution authorizes the practitioner to dispense or prescribe under its registration and assigns a specific internal code number for each practitioner so authorized (See example of a specific internal code number below):



A current list of internal codes and the corresponding individual practitioners is to be maintained by the hospital or other institution. This list is to be made available at all times to other registrants and law enforcement agencies upon request for the purpose of verifying the authority of the prescribing individual practitioner.

Inappropriate Use of the DEA Registration Number

DEA strongly opposes the use of a DEA registration number for any purpose other than the one for which it was intended, to provide certification of DEA registration in transactions involving controlled substances. The use of DEA registration numbers as an identification number is not an appropriate use and could lead to a weakening of the registration system.

The Centers for Medicare and Medicaid Services has developed a National Provider Identification (NPI) number unique to each healthcare provider. The Final Rule for establishment of the NPI system was published in the Federal Register (FR 3434, Vol. 69, No. 15) by the Department of Health and Human Services on January 23, 2004. The effective date of this Final Rule was May 23, 2005; all covered entities must begin using the NPI in standard transactions by May 23, 2007.

Exemption of Federal Government Practitioners from Registration

The requirement of registration is waived for any official of the U.S. Army, Navy, Marine Corps, Air Force, Coast Guard, Public Health Service, or Bureau of Prisons who is authorized to prescribe, dispense, or administer, but not to procure or purchase controlled substances in the course of his/her official duties. Such officials shall follow procedures set forth in Title 21, CFR § 1306 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/her Social Security identification number.

If Federal Government practitioners wish to maintain a DEA registration for a private practice, which would include prescribing for private patients, they must be fully licensed to handle controlled substances by the state in which they are located. Under these circumstances, the Federal Government practitioner will not be eligible for the fee exemption and must pay a fee for the registration.

SECTION III – SECURITY REQUIREMENTS

Required Controls

Title 21, CFR Section 1301.71(a), requires that all registrants provide effective controls and procedures to guard against theft and diversion of controlled substances. A list of factors is used to determine the adequacy of these security controls. Factors affecting practitioners include:

- 1. The location of the premises and the relationship such location bears on security needs
- 2. The type of building and office construction
- 3. The type and quantity of controlled substances stored on the premises
- 4. The type of storage medium (safe, vault, or steel cabinet)
- 5. The control of public access to the facility
- 6. The adequacy of registrant's monitoring system (alarms and detection systems)
- 7. The availability of local police protection

Practitioners are required to store stocks of Schedule II through V controlled substances in a securely locked, substantially constructed cabinet. Practitioners authorized to possess carfentanil, etorphine hydrochloride and/or diprenorphine, must store these controlled substances in a safe or steel cabinet equivalent to a U.S. Government Class V security container.

Registrants should not employ as an agent or employee who has access to controlled substances:

- 1. Any person who has been convicted of a felony offense related to controlled substances
- 2. Any person who has been denied a DEA registration
- 3. Any person who has had a DEA registration revoked
- 4. Any person who has surrendered a DEA registration for cause

Lastly, practitioners should notify the DEA, upon discovery, of any thefts or significant losses of controlled substances and complete a DEA Form 106 regarding such theft or loss.

Safeguards for Prescribers

In addition to the required security controls, practitioners can utilize additional measures to ensure security. These include:

- 1. Keep all prescription blanks in a safe place where they cannot be stolen; minimize the number of prescription pads in use.
- 2. Write out the actual amount prescribed in addition to giving a number to discourage alterations of the prescription order.
- 3. Use prescription blanks only for writing a prescription order and not for notes.
- 4. Never sign prescription blanks in advance.
- 5. Assist the pharmacist when they telephone to verify information about a prescription order; a corresponding responsibility rests with the pharmacist who dispenses the prescription order to ensure the accuracy of the prescription.
- 6. Contact the nearest DEA field office (see Appendix E) to obtain or to furnish information regarding suspicious prescription activities.
- 7. Use tamper-resistant prescription pads.

SECTION IV – RECORDKEEPING REQUIREMENTS

Recordkeeping Requirements

Each practitioner must maintain inventories and records of controlled substances listed in Schedules I and II separately from all other records maintained by the registrant. Likewise, inventories and records of controlled substances in Schedules III, IV, and V must be maintained separately or in such a form that they are readily retrievable from the ordinary business records of the practitioner. All records related to controlled substances must be maintained and be available for inspection for a minimum of two years.

A registered practitioner is required to keep records of controlled substances that are dispensed to the patient, other than by prescribing or administering, in the lawful course of professional practice. A registered practitioner is not required to keep records of controlled substances that are prescribed in the lawful course of professional practice, unless such substances are prescribed in the course of maintenance or detoxification treatment. A registered practitioner is not required to keep records of controlled substances that are administered in the lawful course of professional practice unless the practitioner regularly engages in the dispensing or administering of controlled substances and charges patients, either separately or together with charges for other professional services, for substances so dispensed or administered. A registered practitioner is also required to keep records of an individual.

Inventory

Each registrant who maintains an inventory of controlled substances must maintain a complete and accurate record of the controlled substances on hand and the date that the inventory was conducted. This record must be in written, typewritten, or printed form and be maintained at the registered location for at least two years from the date that the inventory was conducted. After an initial inventory is taken, the registrant shall take a new inventory of all controlled substances on hand at least every two years.

Each inventory must contain the following information:

- 1. Whether the inventory was taken at the beginning or close of business
- 2. Names of controlled substances
- 3. Each finished form of the substances (e.g., 100 milligram tablet)
- 4. The number of dosage units of each finished form in the commercial container (e.g., 100 tablet bottle)
- 5. The number of commercial containers of each finished form (e.g., four 100 tablet bottles)

6. Disposition of the controlled substances

It is important to note that inventory requirements extend to controlled substance samples provided to practitioners by pharmaceutical companies.

Disposal of Controlled Substances

A practitioner may dispose of out-of-date, damaged, or otherwise unusable or unwanted controlled substances, including samples, by transferring them to a registrant who is authorized to receive such materials. These registrants are referred to as "Reverse Distributors." The practitioner should contact the local DEA field office (See Appendix E) for a list of authorized Reverse Distributors. Schedule I and II controlled substances should be transferred via the DEA Form 222, while Schedule III–V compounds may be transferred via invoice. The practitioner should maintain copies of the records documenting the transfer and disposal of controlled substances for a period of two years.

SECTION V – VALID PRESCRIPTION REQUIREMENTS

Prescription Requirements

A prescription is an order for medication which is dispensed to or for an ultimate user. A prescription is not an order for medication which is dispensed for immediate administration to the ultimate user (for example, an order to dispense a drug to an inpatient for immediate administration in a hospital is not a prescription).

A prescription for a controlled substance must be dated and signed on the date when issued. The prescription must include the patient's full name and address, and the practitioner's full name, address, and DEA registration number. The prescription must also include:

- 1. drug name
- 2. strength
- 3. dosage form
- 4. quantity prescribed
- 5. directions for use
- 6. number of refills (if any) authorized

A prescription for a controlled substance must be written in ink or indelible pencil or typewritten and must be manually signed by the practitioner on the date when issued. An individual (secretary or nurse) may be designated by the practitioner to prepare prescriptions for the practitioner's signature.

The practitioner is responsible for ensuring that the prescription conforms to all requirements of the law and regulations, both federal and state.

Who May Issue

A prescription for a controlled substance may only be issued by a physician, dentist, podiatrist, veterinarian, mid-level practitioner, or other registered practitioner who is:

- 1. Authorized to prescribe controlled substances by the jurisdiction in which the practitioner is licensed to practice
- 2. Registered with DEA or exempted from registration (that is, Public Health Service, Federal Bureau of Prisons, or military practitioners)
- 3. An agent or employee of a hospital or other institution acting in the normal course of business or employment under the registration of the hospital or other institution which is registered in lieu of the individual practitioner being registered provided that additional requirements as set forth in the CFR are met.

Purpose of Issue

To be valid, a prescription for a controlled substance must be issued for a legitimate medical purpose by a practitioner acting in the usual course of professional practice. The practitioner is responsible for the proper prescribing and dispensing of controlled substances. In addition, 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 valid prescription within the meaning and intent of the Controlled Substances Act 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.

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.

Schedule II Substances

Schedule II controlled substances require a written prescription which must be signed by the practitioner. There is no federal time limit within which a Schedule II prescription must be filled after being signed by the practitioner.

While some states and many insurance carriers limit the quantity of controlled substance dispensed to a 30-day supply, there are no specific federal limits to quantities of drugs dispensed via a prescription. For Schedule II controlled substances, an oral order is only permitted in an emergency situation.

Refills

The refilling of a prescription for a controlled substance listed in Schedule II is prohibited (Title 21 U.S. Code § 829(a)).

Issuance of Multiple Prescriptions for Schedule II Substances

DEA has revised its regulations regarding the issuance of multiple prescriptions for schedule II controlled substances. Under the new regulation, which became effective December 19, 2007, an individual practitioner may issue multiple prescriptions authorizing the patient to receive a total of up to a 90-day supply of a schedule II controlled substance provided the following conditions are met:

1. Each separate prescription is issued for a legitimate medical purpose by an individual practitioner acting in the usual course of professional practice.

2. The individual practitioner provides written instructions on each prescription (other than the first prescription, if the prescribing practitioner intends for that prescription to be filled immediately) indicating the earliest date on which a pharmacy may fill each prescription.

- 3. The individual practitioner concludes that providing the patient with multiple prescriptions in this manner does not create an undue risk of diversion or abuse.
- 4. The issuance of multiple prescriptions is permissible under applicable state laws.
- 5. The individual practitioner complies fully with all other applicable requirements under the Controlled Substances Act and Code of Federal Regulations, as well as any additional requirements under state law.

It should be noted that the implementation of this change in the regulation should not be construed as encouraging individual practitioners to issue multiple prescriptions or to see their patients only once every 90 days when prescribing schedule II controlled substances. Rather, individual practitioners must determine on their own, based on sound medical judgment, and in accordance with established medical standards, whether it is appropriate to issue multiple prescriptions and how often to see their patients when doing so.

Facsimile Prescriptions for Schedule II Controlled Substances

In order to expedite the filling of a prescription, a prescriber may transmit a Schedule II prescription to the pharmacy by facsimile. The original Schedule II prescription must be presented to the pharmacist for review prior to the actual dispensing of the controlled substance.

In an emergency, a practitioner may call-in a prescription for a Schedule II controlled substance by telephone to the pharmacy, and the pharmacist may dispense the prescription provided that the quantity prescribed and dispensed is limited to the amount adequate to treat the patient during the emergency period. The prescribing practitioner must provide a written and signed prescription to the pharmacist within seven days. Further, the pharmacist must notify DEA if the prescription is not received.

Exceptions for Schedule II Facsimile Prescriptions

DEA has granted three exceptions to the facsimile prescription requirements for Schedule II controlled substances. The facsimile of a Schedule II prescription may serve as the original prescription as follows:

- 1. A practitioner prescribing Schedule II narcotic controlled substances to be compounded for the direct administration to a patient by parenteral, intravenous, intramuscular, subcutaneous or intraspinal infusion may transmit the prescription by facsimile. The pharmacy will consider the facsimile prescription a "written prescription" and no further prescription verification is required. All normal requirements of a legal prescription must be followed.
- 2. Practitioners prescribing Schedule II controlled substances for residents of Long Term Care Facilities (LTCF) may transmit a prescription by facsimile to the dispensing pharmacy. The practitioner's agent may also transmit the prescription to the pharmacy. The facsimile prescription serves as the original written prescription for the pharmacy.
- 3. A practitioner prescribing a Schedule II narcotic controlled substance for a patient enrolled in a hospice care program certified and/or paid for by Medicare under Title XVIII or a hospice program which is licensed by the state may transmit a prescription to the dispensing pharmacy by facsimile. The practitioner or the practitioner's agent may transmit the prescription to the pharmacy. The practitioner or agent will note on the prescription that it is for a hospice patient. The facsimile serves as the original written prescription.

Schedule III-V Substances

A prescription for controlled substances in Schedules III, IV, and V issued by a practitioner, may be communicated either orally, in writing, or by facsimile to the pharmacist, and may be refilled if so authorized on the prescription or by call-in.

Refills

Schedule III and IV controlled substances may be refilled if authorized on the prescription. However, the prescription may only be refilled up to five times within six months after the date on which the prescription was issued. After five refills or after six months, whichever occurs first, a new prescription is required.

Facsimile Prescriptions for Schedule III-V Substances

Prescriptions for Schedules III-V controlled substances may be transmitted by facsimile from the practitioner or an employee or agent of the individual practitioner to the dispensing pharmacy. The facsimile is considered to be equivalent to an original prescription.

Telephone Authorization for Schedule III-V Prescriptions

A pharmacist may dispense a controlled substance listed in Schedule III, IV, or V pursuant to an oral prescription made by an individual practitioner and promptly reduced to writing by the pharmacist containing all information required for a valid prescription, except for the signature of the practitioner.

Delivery of a Controlled Substance to Persons Outside the U.S.

Controlled substances that are dispensed pursuant to a legitimate prescription may not be delivered or shipped to individuals in another country. Any such delivery or shipment is a prohibited export under the CSA.

SECTION VI – OPIOID (NARCOTIC) ADDICTION TREATMENT PROGRAMS

The Narcotic Addiction Treatment Act of 1974 and the Drug Addiction Treatment Act of 2000 amended the CSA with respect to the use of controlled substances in the medical treatment of addiction. These laws established the procedures for approval and licensing of practitioners involved in the treatment of opioid addiction as well as improving the quality and delivery of that treatment to the segment of society in need.

Practitioners wishing to administer and dispense approved Schedule II controlled substances (that is, methadone) for maintenance and detoxification treatment must obtain a separate DEA registration as a Narcotic Treatment Program. Application for registration as a Narcotic Treatment Program. Application to obtaining this separate DEA registration, this type of activity also requires the approval and registration of the Center for Substance Abuse Treatment (CSAT) within the Substance Abuse and Mental Health Services Administration (SAMHSA) of the Department of Health and Human Services (HHS), as well as the applicable state methadone authority.

If a practitioner wishes to prescribe, administer, or dispense Schedule III, IV, or V controlled substances approved for addiction treatment (i.e., buprenorphine drug products), the practitioner must request a waiver (Form SMA-167) and fulfill the requirements of CSAT. CSAT will then notify DEA of all waiver requests. DEA will review each request. If DEA approves this waiver, the practitioner will receive a Unique Identification Number. If a practitioner chooses to dispense controlled substances, the practitioner must maintain, separate from all other records, for a period of at least two years, all required records of receipt, storage, and distribution. If a practitioner chooses to prescribe these controlled substances, the practitioner must utilize their Unique Identification Number on the prescription in addition to his/her regular DEA registration number. The practitioner must also maintain a record of each such prescription for a period of at least two years. Practitioners should be aware that there may be limits on how many patients they may treat for opioid addiction at any given time and should check with SAMHSA to determine these limits.

Note that not all treatment programs utilize controlled substances, that is, some are drug free. Accordingly, these activities do not require DEA registration or approval.

Practitioners can find additional information regarding addiction treatment by visiting DEA's Office of Diversion Control website at www.DEAdiversion.usdoj.gov. Click on "Publications," then "Narcotic Treatment Programs: Best Practices Guidelines." The DEA application Form 363 may be completed on-line.

To learn more about CSAT's requirements, practitioners may visit one or more of the following websites: <u>www.samhsa.gov/centers/csat2002/csat_frame.html</u>, <u>www.csat.samhsa.gov</u>, or <u>www.buprenorphine.samhsa.gov</u>.

If the practitioner has a patient who is in need of addiction treatment, but does not wish to treat the individual, the practitioner can refer the patient to an existing facility through the following website: <u>www.findtreatment.samhsa.gov</u>.

APPENDICES

APPENDIX A

CSA & CFR Definitions

Administer

The direct application of a controlled substance to the body of a patient or research subject by 1) a practitioner or (in his presence) by his authorized agent, or 2) the patient or research subject at the direction and in the presence of the practitioner, whether such application is by injection, inhalation, ingestion, or any other means.

Dispense

To deliver a controlled substance to an ultimate user or research subject by, or pursuant to the lawful order of, a practitioner, including the prescribing and administering of a controlled substance and the packaging, labeling, or compounding necessary to prepare the substance for such delivery.

Dispenser

An individual practitioner, institutional practitioner, pharmacy or, pharmacist who dispenses a controlled substance.

Individual Practitioner

A physician, dentist, veterinarian, or other individual licensed, registered or otherwise permitted, by the United States or the jurisdiction in which they practice, to dispense a controlled substance in the course of professional practice, but does not include a pharmacist, a pharmacy, or an institutional practitioner.

Institutional Practitioner

A hospital or other person (other than an individual) licensed, registered or otherwise permitted, by the United States or the jurisdiction in which it practices, to dispense a controlled substance in the course of professional practice, but does not include a pharmacy.

Inventory

All factory and branch stocks in finished form of a basic class of controlled substance manufactured or otherwise acquired by a registrant, whether in bulk, commercial containers, or contained in pharmaceutical preparations in the possession of the registrant (including stocks held by the registrant under separate registration as a manufacturer, importer, exporter, or distributor).

Long Term Care Facility

A nursing home, retirement care, mental care, or other facility or institution which provides extended health care to resident patients.

Mid-level Practitioner

An individual practitioner, other than a physician, dentist, veterinarian, or podiatrist, who is licensed, registered or otherwise permitted by the United States or the jurisdiction in which he/she practices, to dispense a controlled substance in the course of professional practice. Examples of mid-level practitioners include, but are not limited to, health care providers such as nurse practitioners, nurse midwives, nurse anesthetists, clinical nurse specialists, and physician assistants who are authorized to dispense controlled substances by the state in which they practice.

Pharmacist

Any pharmacist licensed by a state to dispense controlled substances, and shall include any other person (e.g., pharmacist intern) authorized by a state to dispense controlled substances under the supervision of a pharmacist licensed by such state.

Prescription

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).

Readily Retrievable

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, redlined, or in some other manner visually identifiable apart from other items appearing on the records.

APPENDIX B

Questions and Answers

The following questions are those that are frequently encountered by DEA's Office of Diversion Control and its field units. These questions and their accompanying answers are provided in context of the CSA and its federal regulations.

${f Q}$ Are separate registrations required for separate locations?

A A separate registration is required for each principal place of business or professional practice where controlled substances are stored or dispensed by a person.

Q Does a practitioner need a separate registration to treat patients at remote health care facilities?

A Separate registration is not required in 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.

${f Q}$ Do all practitioners in a group practice need to be registered?

A An individual practitioner who is an agent or employee of another practitioner (other than a mid-level practitioner) registered to dispense controlled substances may, when acting in the normal course of business or employment, administer or 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 or she practices, under the registration of the employer or principal practitioner in lieu of being registered him/herself.

${f Q}$ Do medical residents assigned to hospitals need to register?

A An individual practitioner who is an agent or employee of a hospital or other institution may, when acting in the normal course of business or employment, administer, dispense, or prescribe controlled substances under the registration of the hospital or other institution which is registered in lieu of being registered provided that additional requirements as set forth in the CFR are met.

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Q Are military personnel exempted from registration?

A Registration is waived for any official of the U.S. Army, Navy, Marine Corps, Air Force, or Coast Guard who is authorized to prescribe, dispense, or administer, but not procure or purchase, controlled substances in the course of his/her official duties. Such officials must follow procedures set forth in 21 CFR Part 1306 regarding prescriptions. Branch of service or agency and the service identification number of the issuing official is required on the prescription form in lieu of the DEA registration number.

If any exempted official engages as a private individual in any activity or group of activities for which registration is required, that individual must obtain a registration for those private activities.

Further, practitioners serving in the U.S. Military are exempt from registering with DEA, but are not authorized to procure or purchase controlled substances in the course of their official duties.

A number of states also require military practitioners to acquire a separate state license if they issue prescriptions that are filled outside the military facility where they practice.

Q Are contract practitioners working at U.S. Military Installations also exempt from registration?

A They are not exempt. A contract practitioner who is not an official of the military on active duty, but is engaged in medical practice at a military installation, must possess a current DEA registration. The individual must also possess a valid state license for the same state in which he/she is registered with DEA.

${f Q}$ What should a practitioner do if he/she discovers a theft or loss?

A Registrants must notify the DEA field office in their area of the theft or significant loss of any controlled substances upon discovery. The registrant must also complete DEA Form 106 documenting the loss or theft.

${f Q}$ What is meant by "acceptable medical practice?"

A The legal standard that a controlled substance may only be prescribed, administered, or dispensed for a legitimate medical purpose by a physician acting in the usual course of professional practice has been construed to mean that the prescription must be "in accordance with a standard of medical practice generally recognized and accepted in the United States."

Federal courts have long recognized that it is not possible to expand on the phrase "legitimate medical purpose in the usual course of professional practice" in a way that will provide definitive guidelines to address all the varied situations physicians may encounter.

While there are no criteria to address every conceivable instance of prescribing, there are recurring patterns that may be indicative of inappropriate prescribing:

- An inordinately large quantity of controlled substances prescribed or large numbers of prescriptions issued compared to other physicians in an area;
- No physical examination was given;
- Warnings to the patient to fill prescriptions at different drug stores;
- Issuing prescriptions knowing that the patient was delivering the drugs to others;
- Issuing prescriptions in exchange for sexual favors or for money;
- Prescribing of controlled drugs at intervals inconsistent with legitimate medical treatment;
- The use of street slang rather than medical terminology for the drugs prescribed; or
- No logical relationship between the drugs prescribed and treatment of the condition allegedly existing.

Each case must be evaluated based on its own merits in view of the totality of circumstances particular to the physician and patient.

For example, what constitutes "an inordinately large quantity of controlled substances," can vary greatly from patient to patient. A particular quantity of a powerful Schedule II opioid might be blatantly excessive for the treatment of a particular patient's mild temporary pain, yet insufficient to treat the severe unremitting pain of a cancer patient.

${f Q}$ What information is required to be provided on a written prescription?

A All written prescriptions for controlled substances must be dated as of, and signed on, the date when issued. Each prescription must indicate the full name and address of the patient, the drug name, strength, dosage form, quantity prescribed,

2006 Edition Page 30 directions for use and the name, address, and DEA number of the practitioner. Further, prescriptions must be written in ink, indelible pencil, or by typewriter, and must be manually signed by the practitioner.

Q What is meant by "date of issuance?"

A The date a prescription is issued is the same date that the prescribing practitioner actually writes and signs the prescription.

${f Q}$ Is there a time limit for filling Schedule II prescriptions?

A There is no federal time limit for filling Schedule II prescriptions. However, some state laws do set time limits.

APPENDIX C

Summary of Controlled Substances Act Requirements

	Schedule II	Schedule III & IV	Schedule V
Registration	Required	Required	Required
Receiving Records	Order Forms (DEA Form-222)	Invoices, Readily Retrievable	Invoices, Readily Retrievable
Prescriptions	Written Prescription (See exceptions*)	Written, Oral, or Fax	Written, Oral, Fax, or Over The Counter**
Refills	No	No more than 5 within 6 months	As authorized when prescription is issued
Distribution Between Registrants	Order Forms (DEA Form-222)	Invoices	Invoices
Security	Locked Cabinet or Other Secure Storage	Locked Cabinet or Other Secure Storage	Locked Cabinet or Other Secure Storage
Theft or Significant Loss	Report and complete DEA Form 106	Report and complete DEA Form 106	Report and complete DEA Form 106

Note: All records must be maintained for 2 years, unless a state requires a longer period.

- * Emergency prescriptions require a signed follow-up prescription.
 Exceptions: A facsimile prescription serves as the original prescription when issued to residents of Long Term Care Facilities, Hospice patients, or compounded IV narcotic medications.
- ** Where authorized by state controlled substances authority.

APPENDIX D

Internet Resources

DEA's Diversion Control Program Website www.DEAdiversion.usdoj.gov

DEA Homepage www.dea.gov

<u>U.S. Government Printing Office</u> www.gpoaccess.gov/cfr/index.html

Provides access to the Code of Federal Regulations (21 CFR, Parts 1300 to end), primary source for the Practitioner's Manual, and the Federal Register which contains proposed and finalized amendments to the CFR.

Office of National Drug Control Policy (ONDCP) www.whitehousedrugpolicy.gov

Food and Drug Administration www.FDA.gov

HHS & SAMHSA's National Clearinghouse for Alcohol and Drug Information www.health.org

<u>SAMHSA/CSAT</u> www.csat.samhsa.gov

Federation of State Medical Boards www.FSMB.org

National Association of Boards of Pharmacy www.nabp.net

National Association of State Controlled Substances Authorities www.nascsa.org

APPENDIX E

Drug Enforcement Administration Diversion Field Office Locations

For address and telephone number updates, please see the DEA website: www.deadiversion.usdoj.gov/offices_n_dirs/index.html

Appendix E pages 34-39 of this manual contained outdated Field Office Information and therefore have been removed. Please refer to the above link for current Diversion Field Office Locations.

APPENDIX F

Small Business and Agriculture Regulatory Enforcement Ombudsman

The Small Business and Agriculture Regulatory Enforcement Ombudsman and 10 Regional Fairness Boards were established to receive comments from small businesses about federal agency enforcement actions. The Ombudsman will annually evaluate the enforcement activities and rate each agency's responsiveness to small business. If you wish to comment on DEA enforcement actions, you may contact the Ombudsman at 1-888-REG-FAIR (1-888-734-3247).

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APPENDIX G

Additional Assistance

This publication is intended to provide guidance and information on the requirements of the Controlled Substances Act and its implementing regulations. If you require additional clarification or assistance, or wish to comment on any matter regarding the DEA's requirements or regulatory activities, please contact your local DEA Diversion field office (see Appendix E). Every effort will be made to respond promptly to your inquiry.

<u>Plain Language</u>

The Drug Enforcement Administration has made every effort to write this manual in clear, plain language. If you have suggestions as to how to improve the clarity of this manual, please contact us at:

Drug Enforcement Administration Office of Diversion Control Liaison and Policy Section Washington, D.C. 20537 Telephone: (202) 307-7297

APPENDIX H – DEA FORMS

The following pages provide samples of several forms frequently encountered by DEA registrants. Included are:

- **DEA Form 41** Registrants Inventory of Drugs Surrendered
- **DEA Form 106** Report of Theft or Loss of Controlled Substances
- DEA Form 222 U.S. Official Order Form for Controlled Substances
- **DEA Form 224** Application for Registration
- DEA Form 224a Renewal Application for DEA Registration
- **DEA Form 363** Application for Registration as a Narcotic Treatment Program
- **DEA Form 363a** Renewal Application for DEA Registration as a Narcotic Treatment Program

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L			L	-	s DEA Number						
_		Registrant's Tel									
NOTE: CERTIFIED M	AIL (Return Receipt Requested) IS REQUIRED FOR SHIPM IA U.S. POSTAL SERVICE. See instructions on reverse (par	IENTS									
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	NAME OF DRUG OR PREPARATION	Number of Con-	grams, tablets, ounces or	Sub- stance	FOR DE/						
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REPORT OF THEFT OR LOSS OF CONTROLLED SUBSTANCES

Federal Regulations require registrants to submit a Enforcement Administration.	detailed report of any theft or los	ss of Controlled Substances	to the Drug	OMB APPROVAL
Complete the front and back of this form in triplicat Retain the triplicate copy for your records. Some st	e. Forward the original and dup ates may also require a copy of	plicate copies to the nearest f this report.	DEA Office.	No. 1117-0001
1. Name and Address of Registrant (include ZIP Code))		2. Phone N	lo. (Include Area Code)
		ZIP CODE		. ,
3. DEA Registration Number	4. Date of Theft or Loss	5. Principal Business of Re	gistrant (Chec	k one)
2 ltr. prefix 7 digit suffix		1 Pharmacy	5 [Distributor
		2 Practitioner	6 [Methadone Program
		3 🗌 Manufacturer	7 [Other (Specify)
		4 🗌 Hospital/Clinic		
6. County in which Registrant is 7. Was Theft rep located to Police?	orted 8. Name and Teleph	one Number of Police Depart	ment (Include	Area Code)
located to Police?				
	- No			
Yes	_ 140			
9. Number of Thefts or Losses Registrant 10. Type of	of Theft or Loss (Check one an	d complete items below as	appropriate)	
has experienced in the past 24 months	linkt brock in a 🗆 a			
		ployee pilferage 5	vv.	
2 🗆 🌶	Armed robbery 4 🗌 Cus	tomer theft 6	Lost in t	ransit (Complete Item 14)
11. If Armed Robbery, was anyone:	 Purchase value to Controlled Substa 	registrant of 13.		armaceuticals or
	Controlled Substa	noes taken?	merchandise	Yes (Est. Value)
Killed? No Yes (How many)	— .		_	
Injured? No Yes (How many)	\$		\$	
14. IF LOST IN TRANSIT, COMPLETE THE FOLLOWI	NG:			
A. Name of Common Carrier E	Name of Consignee	C. (Consignee's D	EA Registration Number
D. Was the carton received by the customer?	. If received, did it appear to b	e tampered with? F. F fi		rienced losses in transit carrier in the past?
Yes No	🗌 Yes 🗌 N	• []No []	Yes (How Many)
15. What identifying marks, symbols, or price codes w	ere on the labels of these conta	ainers that would assist in i	dentifying the	products?

16. If Official Controlled Substance Order Forms (DEA-222) were stolen, give numbers.

17. What security measures have been taken to prevent future thefts or losses?

AUTHORITY: Section 301 of the Controlled Substances Act of 1970 (PL 91-513). PURPOSE: Report theft or loss of Controlled Substances. ROUTINE USES: The Controlled Substances Act authorizes the production of special reports required for statistical and analytical purposes. Disclosures of the information from this system are made to the following categories of users for the	accordance with the Paperwork Reduction Act of 1995, no person is uired to respond to a collection of information unless it displays a ly d OMB control number. The valid OMB control number for this ection of information is 1117-0001. Public reporting burden for this ection of information is estimated to average 30 minutes per ponse, including the time for reviewing instructions, searching sting data sources, gathering and maintaining the data needed, and spletting and reviewing the collection of information.

FORM DEA - 106 (11-00) Previous editions obsolete

CONTINUE ON REVERSE

Trade Nan	ne of Substance or Preparation	Name of Controlled Substance in Preparation	Dosage Strength and Form	Quantity
Examples:	Desoxyn	Methamphetamine Hydrochloride	5 mg Tablets	3 x 100
	Demerol	Meperidine Hydrochloride	50 mg/ml Vial	5 x 30 ml
	Robitussin A-C	Codeine Phosphate	2 mg/cc Liquid	12 Pints
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9. 0.				
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Signature

Date

DEPICTION of PAGE 1 of DEA FORM-222 U.S. OFFICIAL ORDER FORM - SCHEDULES I & II

Se		of PURCHAS		m may be issued for Schedule I and II substances unless a pplication form has been received, (21 CFR 1305.04). OMB APPRO No. 1117-001														
TO:	(Name of Su	pplier)				STREET	AD	DRE	ESS									
CIT	Y and STAT			DATE			TO BE FILLED IN BY SUPPLIER											
							SL	JPP	LIEF	RS D	ΕA	REG	SIST	rr/	ATIC	N N	No.	
L		TO BE FIL		/ PURCHASI	ER													
I N E No.	No. of Size of Packages Package			Name of	National Drug Code										Packages Shipped	Date Shipped		
1																T		
2																		
3																		
4																		
5																		
6																		
7																		
8																T		
9																		
10																		
		LAST LINE COMPLETED) (MUST	T BE 10 OR LI	ESS)	SIGNATURI OR ATTORI												
Dat	e Issued		DEA Regis	stration No.	Nar	me and Addre	ess	of R	egis	tran	t							
Sch	edules		<u> </u>															
Reç	jistered as a		No. of this	Order Form	n I													
	A Form-222 t. 1992)		U			ORDER F INFORCEME SUPPLIE	NT	ADI	MIN	STF			.ES	518	& II			

Note: The graphic illustrated above is not intended to be used as an actual order form.

Form-224	APPLICATION FOR REGISTRATION Under the Controlled Substances Act	APPROVED OMB NO 1117-00 FORM DEA-224 (9- Previous editions are obsol
NATOLATIONS		REGISTRATION INFORMATION :
INSTRUCTIONS	 To apply by mail complete this application. Keep a copy for your records. Print clearly, using black or blue hk, or use a typewriter. Mail this form to the address provided in Section 7 or use enclosed envelope. Include the correct payment amount. FEE IS NON-REFUNDABLE. If you have any questions call 600-682-6930 prior to submitting your application. Save time - apply online at www.deadlwerston.usdol.gov. 	
	IMPORTANT: DO NOT SEND THIS APPLICATION AND APPLY ONLINE.	
		* ~~~~~~
		\$390.00
APPI	JCANT	FEE IS NON-REFUNDABLE
SECTION I IDEN	TIFICATION	
Last Name (if regist	ration is for individual) -OR- Business or Facility Name (if registration is for bus	ness entry)
First Name (if regist	ration is for individual)	Midd
Business or Facility	Name 2 ("doing business as", continuation of business name, or name of fee exempt ins	titution)
Address Line 1 (stre	et address)	
Address Line 2		
City		State Zip Code
Durchase Dhana hir	mbar Durkers Fau Munker	
Business Phone Nu	mber Business Fax Number	
INFORMATION	Tax Identification Number (I'registration is for business) Social Se	Curity Number (if registration is for individual) Provide SSN or Ti
Mandatory pursuant to Debt Collection Improvements Act		See note #3 on bottom of page 2
SECTION 2	Hospital/Clinic Ambulance Service Practiti	MD, DO, DPM, DVM, MD or PHD) DEGREE
BUSINESS ACTIVITY Check one box only	Nursing Home Animal Shefter Practiti	MD, DO, DPM, DVM, MD or PHD) Practitioners and MLF Enter your profession dographic fragmitst
See page 3 for additional instructions	Control Ell Diservoy	el Practitioner (MLP) MD, MP, ND, NP, OD, PA, or RPH)
I	Retail Pharmacy Automated Dispensing System Euthan	asia Technician
FOR Automated Dispensing (ADS) ONLY:	System DEA Registration # of Retail Pharmacy for this ADS	An ADS is automatically lee-exempt. Skip Section 6 and Section 7 on page 2. You must attach a notorized attidavil.
SECTION 3	Schedule II Narcotic Schedule III Narcotic	Schedule IV
DRUG SCHEDULES Check all that apply	Schedule II Non-Narcotic Schedule III Non-Narcotic	Schedule V
	Check this box if you require official order forms for purchase of schedule II narcotic/schedule II non-narcotic controlled substances	

SECTION 4	the sch	edules főr w	hich you a	o prescr re apply	ibe, di /ing un	stribute ider the	e, disp e laws	ense, o of the	state o	x rese or juri:	sard sdic	h, or c tion in	whic	wise i sh you	hand u are	e the controlled substa operating or propose to	nces in o operate?
STATE LICENSE(S) Be sure to include both	YES	PENDING	NO						Π		Т	Т	Т	Π		State License Number	
state license numbers li applicable	1	H	H.	Ħ	Ħ	-	Ħ		Ħ	-	÷	÷	÷	Ħ	H	State Controlled Su	
											_	_	_			License Number (if	
SECTION 5 LIABILITY	1. Hasth	ə applicant e	ver been c	onvicte	ed of a	crime	in cor	nnectio	n with	contr	ollec	d subs	stance	e(s) u	under	r state or federal law?	YES NO
IMPORTANT	restrict	ed, or denied	1?			,										woked, suspended,	
All questions in this section must be answered.	3. Has the revoke	e applicant e d, suspende	d, denied, i	dered (f restricte	or cau d, or p	se) or f laced (had a on pro	state p bation	rofess ? Is ar	ional ny su	licer ch a	nse or action	pen	ding	1 sub ?	stance registration	
	control registra	e applicant is a corporation (other than a corporation whose stock is owned and traded by the public), association, thership, or pharmacy, has any officer, partner, stockholder, or proprietor been convicted of a crime in connection with invited substance(s) lunder state or federal law, or ever summendered, for cause, or had a federal controlled substance istration revoked, suspended, restricted, denied, or ever had a state professional license or controlled substance istration revoked, suspended, denied, restricted or placed on probation?															
EXPLANATION OF "YES" ANSWERS	Date(s)	s) of Incident: Location(s) of Incident:															
Applicants who have answered "YES" to any of the four question above must provide a statement to explain such answers	nava Nature of Incident: to uestions vide axplain																
Use this space or altack a separate sheet and return with application	^h Result	of incident:															
SECTION 6	Check this box if the applicant is a federal, state, or local government operated hospital, institution or official. Be sure to enter the name and address of the exempt institution in Section 1.																
CERTIFICATION OF EXEMPTION from application fee	The un Instituti		ereby certif	les that	the ap	plicant	name	xi here	on is a	a fede			or loc	ai go	wern	ment-operated hospital	
Provide the name and phone number of the certifying official	Signati	ire of certify	ng official (other th	an app	llcant)							_	Date	•		
	Print or	type name	and title of	cərtifyin	ng offic	ial								Tele	phon	e No. (required for verification	stion)
SECTION 7 METHOD OF	Ch Ch	Make Sck Seep	check paya age 4 of ins	ble to: D tructions	rug En for imp	forcem ortant in	ent Ad Normal	ministr ion.	ation								
PAYMENT Check one form of	🗌 Am	erican Expre	66 🗌 D	scover		Master	Card		Visa							Mail this form with pa	
payment only	Credit	Credit Card Number Expiration Date									_	U.S. Department of Justice Drug Enforcement Administration					
						-	Ц	-			-	1.				P.O. Box 280 Washington, DC 20	
Sign If paying by credit card	Signati	ure of Card H	lokler													FEE IS NON-REFU	NDABLE
	Printed	Name of Ca	ard Holder														
SECTION 8	I certify	that the fore	egoing info	rmation	fumisi	hed on	this a	pplicati	ion ist	rue ar	nd c	orrect					
APPLICANT'S SIGNATURE Sign in Ink	Signat	ure of appli	cant											ī	Date		
	Print or	type name	and title of	applica	nt												
	WARNII	NG: Section & nt information	43(a)(4)(A) o In the applic	f Title 21 ation is s	, United subject i	d States to Impri:	Code	states ti nt for no	hat any st more	perso than fe	n wh our y	o know ears, a	vingly 1 fine	or ink of not	more	ally furnishes faise or than \$30,000, or both.	
valid OMB control nu the time for reviewing 3. The Debt Collection This number is requir 4. PRIVACY ACT INFO	he Paperwo Imber for the Improveme red for debt RMATION	rk Reduction A is collection is is, searching e nts Act of 1996 collection pro	Act of 1995, i 1117-0014, xisting data 5 (PL 104-13 cedures sho	no perso Public n sources, 4) requir uld your	n is req eporting gather es that fee bec	uired to burden ing and you furr come un	respon for the mainta hish you collect:	nd to a c s collect lining th ur Taxpa able.	collectio tion of it e data i ayer i de	in of in nforma needer artifyin	stion d, an ig Nu	is estin d comp imber a	mated pietin; and/oi	t to av g and r Sock	erage revier al Sec	a valid OMB control numb a 12 minutes par response, wing the collection of inform suffy Number on this appli	including nation. cation.
AUTHORIT PURPOSE: ROUTINE U EFFECT:	Y:	texpayer iden To obtain info The Controlle Information fro A. Other feder B. State and I	tilying numb rmation requid Substance om this syste rai law enfor ocal law enfor gistered und	er and/or ired to re is Act Re or are m cement a proement ler the C	r social agister a gistratic ade to t and regi t and re ontrolle	security applican on Reco the follo ulatory z guiatory d Subst	numbe Its purs rds pro wing ca sgencia agencia ances /	ar). Laint to iduces s ategoria is for lay lies for l Act (PL	the Cor special is of use w enfor law enfo 91-513	reports ars for cemen	d Sul s as the p at ani ent a	bstance require purpos d regula ind regula	es Act ed for : es sta latory ulator	t of 19 statist ated: purpo ry purp	170. lical a xes. poses	nenks Act of 1998 (PL 104 natylical purposes. Disclor gistration of customers.	
erest.			-price form v	predu	and prot		EW - P										

Form-224	APPLICATIO	ON FOR REGIS	TRATION								
	Supplementary	Instructions and In	formation								
ADDITIONAL INSTRUCTIONS	SECTION 1. APPLICANT IDENTIFICATION - Information must be typed or printed in the blocks provided to help reduce data entry errors. Fee exempt applications must list the name and address of the fee exempt institution. A physical address is required; after the street address a post office box may be included. Applicant must enter a valid social security number (ISSN), or a tax identification number (TIN) if applying as a business entity. Debt collection information is mandatory pursuant to the Debt Collection improvement Act of 1996.										
	SECTION 2. BUSINE	SS ACTIVITY - Indicate or	ly one. Practitioners al	so enter one degree from t toices: DOM, HMD, MP, N	Nalist DDS, DMD, DO	, DPM, DVM, MD or PHD.					
	Affidavit mústinclude 3) Permit or license nu 4) Required Statemen	 Name of parent retail ph imber(s) and date issued of tribs affidavit is submitte commence proceedings material information con corporation/partnershipt 	armacy and complete a of State certification to or of to obtain a DEA regis to deny the application tained in this all davit in business to prosecution	nacy and attach a notorize ddress 2) Name of Long-te perate ADS at named LTC fation number. If any math under section 304 of the A sy subject the person sign under section 403 of the A	Im Care (LTC) facility : facility vial information is false, ct (21 U.S.C. 6224(a)). ing this affidevit, and the ct (21 U.S.C 643).	and complete address , the Administrator may Any false or fraudulent e named					
	 Name of corporation operating the retail pharmacy 6) Name and title of corporate officer signing affidavit 7) Signafure of authorized officer SECTION 3. DRUG SCHEDULES - Applicants should check all drug schedules to be handled. However, applicants must etil comply with state requirements; federal registration does not overrule state restrictions. Check the order form box only if you intend to purchase or to transfer schedule il controlled substances. Order forms will be matted to the registrated address following issuance of a Cartificate of Registration. SECTION 4. STATE LICENSE(8) - Federal registration by DEA is based upon the applicant's compliance with applicable state and local laws. Applicants soluid onticat the local state localistate local state local is application. If you state requires a separate controlled substance number, provide that number on this application. If a state license has not yet been issued, Indicate "Pending". If state locanis must answer all four questions for the application to be accepted for processing. If you answered "Yes" to any question, provide an explanation in the space provided. If additional space is required, you may attach a separate sheet of paper. SECTION 6. CERTIFICATE OF EXEMPTION - Exemption from payment of application as limited to federal, state or local government operated hospitals, institutions and officials. The applicant's superior reagency officer must certify exempt status. The signature, authority the, and telephone number of the certifying official (other than the application) the provided. 										
	SECTION 7. METHO	D OF PAYMENT - Indicate	the desired method of ;	payment. Make checks par not be accepted. FEES AR	vable to "Drug Enforcen	nent Administration".					
	SECTION 6. APPLIC	ANT'S SIGNATURE - Musi	t be the original signatur	e (in ink) of the applicant.							
CONTACT INFORMATION	ATLANTA DIVISION (ATTN: Registration 75 Spring Street, SW, Atlanta, GA 30303		DETROIT DIVISION (431 Howard Street Detroit, MI 45226		PHILADELPHIA DIVISION OFFICE William J. Green Federal Building 900 Arch Street, Room 10224 Philadelphia, PA 19105						
1. INTERNET www.deadlversion.usdoj.gov	Georgia North Carolina South Carolina	(000) 000-9935 (000) 219-0609	Kentucky Michigan Ohio	(800) 230-6844 (800) 230-6844 (800) 230-6844	Delaware Pennsylvania	(600) 393-8231 (600) 393-8231					
2. TELEPHONE Headquarters Call Center	Tennessee BOSTON DIVISION O JFK Federal Building	(066) 533-6903 (000) 219-7095 (FFICE	EL PASO DIVISION C El Paso Federal Justik 600 South Mesa Hills El Paso, TX 79912	ce Center	PHOENIX DIVISION OFFICE 3010 N. 2nd Street, Suite 301 Phoenix, AZ 55012						
(800) 882-9539	15 New Sudbury Stree Boston, MA 02203-013		New Mexico	(915) 832-6014	Arizona	(800) 741-0902					
3. WRITTEN INQUIRIES DEA P.O. Box 26063	Connecticut Maine Massachusetts	(617) 557-2200 (660) 272-5174 (617) 557-2465	HOUSTON DIVISION 1433 West Loop Sout Houston, TX 77027-9	h. Sulte 600	SAN DIEGO DIVISION OFFICE 4550 Viewrldge Avenue San Diego, CA 02123-1637						
Washington DC 20038-8083 4. DEA OFFICES	New Hampshire Rhode Island	(000) 272-5174 (617) 557-2200	Texas (S. & Central)		California (Southern)						
DEA Offices are listed (600, 677, and 665	CARIBBEAN DIVISIO P.O. Box 2167		LOS ANGELES DIVI: 255 East Temple Stre Los Angeles, CA 9001	et, 20th Floor	SAN FRANCISCO DIVISION OFFICE 450 Golden Gate Avenue, 14th Floor R.O. Box 38035 San Francisco, CA 94102						
are toll-free numbers)	San Juan, PR 00922-3 Puerto Rico	(707) 775-1786	California (S. Central) Hawali	(666) 415-9622	California (Northern)						
	U.S. Virgin Islands CHICAGO DIVISION	(787) 775-1786 OFFICE	Nevada Trust Territory	(855) 415-9522 (213) 894-2216	SEATTLE DIVISION (400 Second Avenue, 1 Seattle, WA 98119						
	Nuczynski Federal Bu 230 S. Dearborn Stree Chicago, IL 60604	liding	MIAMI DIVISION OFF 6400 N.W. 53rd Street Miami, FL 33166		Alaska Idaho Oregon	(600) 219-4261 (600) 219-4261 (600) 219-4261					
	Illinois Indiana	(312) 353-1234 (312) 353-1236	Florida	(305) 590-4880	Washington	(000) 219-1410					
	Minnesota North Dakota Wisconsin	(312) 353-9166 (312) 353-9166 (312) 353-1236	NEWARK DIVISION (60 Mulberry Street, 2r Newark, NJ 07102		ST. LOUIS DIVISION 317 South 16th Street St. Louis, MO 63103						
	DALLAS DIVISION O 10160 Technology Blv Dalas, TX 75220		New Jersey NEW ORLEANS DIVI	(888) 356-1071 SION OFFICE	lowa Kansas Missouri	(688) 803-1179 (688) 803-1179 (688) 803-1179					
	Oklahoma Texas (Northern)	(000) 336-4704 (000) 336-4704	3636 N. Causeway Bi Lakeway III, Sulle 160 Metairle, LA 70002	vd	Nebraska South Dakota	(688) 803-1179 (688) 803-1179					
	DENVER DIVISION O 115 Inverness Drive, E Englewood, CO 60112	ast	Alabama Arkansas Louisiana Mississippi	(658) 514-5051 (658) 514-7302 (658) 514-7302 (658) 514-7302	WASHINGTON, D.C. DIVISION OFFICE Techworld Plaza 600 K. Street, N.W., Sulle 500 Washington, D.C. 20001						
	Colorado Montana Utah Wyoming	(800) 326-6900 (800) 326-6900 (800) 326-6900 (800) 326-6900	NEW YORK DIVISION 90 Tenth Avenue New York, NY 10011		District of Columbia Maryland Virginia West Virginia	(677) 601-7974 (677) 330-6670 (677) 601-7974 (677) 330-6670					
NEW INST - Page 3			New York	(677) 663-5769 (212) 337-1593 (212) 337-1594							

DRUG		ned drug code n	umbers. If you are in need of additional information, see 21 0	FR 1306
SCHEDULES	or contact the DEA office serving your area.	-		
	SCHEDULEI		SCHEDULE III	
	NARCOTIC & NON-NARCOTIC		NARCOTIC BASIC CLASSES	CODE
	BASIC CLASSES	CODE	Buprenorphine	9064
	Acetophine	9319	Codeline up to 90 mg/du plus other ingredients	9319
	Acetytmethadol Allytprodine	9601	Dihydrocodeineup to 90 mg/du plus other ingredients Ethylmorphine up to 15 mg/du plus other ingredients	9607
	Alphacetylmethadol (except LAAM)	9603	Hydrocodone up to 15 mg/du plus other ingredients	9006
	Burotenine	7433	Morphine up to 50 mg/100mi or am plus other ingred.	9610
	Dextromoramide	9613	Oplum up to 500 mg/100m, plus other active ingred.	9009
	Diethyttryptamine (DET) 2,5 - Dimethoxyamphetamine (DMA)	7434 7396	NON-NARCOTIC BASIC CLASSES	CODE
	Dimethytiryptamine (DMT)	7435		CODE
	Etorphine (except hydrochloride salt)	9056	Anabolic Steroids	4000
	gamima-Hýdroxýbutýric acid (exceptídrug product)	2010	Berzphetamine	1228
	Heroin Ibogaine	9200 7260	Butaibitai Dronabinoi Pharmaceuticai Product	2100
	Ketobernidone	9625	GHB Drug Product (gamma-Hydroxybutyric acid)	2010
	Lysergic acid diethytamide (LSD)	7315	Ketamine	7265
	Marihuana Mescaline	7360 7361	Methypryton Exclobational plus accordinated active interactions	2575 2271
	Methaguaione	2565	Penióbarbital plus noncontrolled active ingredients Peniobarbital suppository	2271
	3,4 - Methylenedioxyamphetamine (MDA)	7400	Phendimetrazine	1615
	3,4 - Methylenedioxymethamphetamine (MDMA)	7405	Secobarbital plus noncontrolled active ingredients	2316
	n- Ethyl - 1 - Phenylcyclohexylamine (PCE) Peyote	7455 7415	Secobarbital suppository Thiopental	2316 2329
	1 - (1-Phenylcyclohexyl)pyrrolidine (PCP)	7455	Vinbarbital	2335
	Psilocybin	7437		
	Psilocyn	7438		
	Tetrahydrocannabinols (THC) 1-[1-(2-Thieny()-cyclohexyl -piperidine	7370	SCHEDULE IV	
	-[-(2-1111)); of a sum of the sum	1410	NARCOTIC BASIC CLASSES	CODE
	SCHEDULE II		Dextropropoxyphene du Difenoxin 1mg/25ug atropine SO4/du	9278 9167
	NARCOTIC BASIC CLASSES	CODE	NON-NARCOTIC BASIC CLASSES	CODE
	Alphaprodine Anlieridine	9010	Alprzolam	2002
	Cocaine	9041	Barbital	2145
	Codeine	9050	Chioral Hydrate	2465
	Dextropropoxyphene (bulk)	9273	Chiordiazepoxide Ciorazepate	2744 2768
	Diphenoxylate Diprenorphine (M50-50)	9170	Diazepam	2765
	Ethylmorphine	9190	Diethýtpropion	1610
	Etorphine Hydrochloride (M-99)	9059	Fenflüramine	1670 2767
	Glufethimide	2550 9193	Flurazepam Halazepam	2762
	Hydromorphone	9150	Lorazepam	2005
	Lévo-alphacetylmethadol (LAAM)	9645	Mazindol	1605
	Levorphanol	9220	Mebutamate Mephobarbital (Methylphenobarbital)	2800 2250
	Meperidine Methadone	9230 9250	Meprobamate	2620
	Morphine	9300	Methohexital	2264
	Oplum, powdered	9639	Midazolam Oxazepam	2664 2635
	Oplum, raw Oxycodone	9800 9143	Paraidehyde	2565
	Oxymorphone	9652	Pempline	1530
	Poppy Straw	9671	Peniazocine	9709
	Poppy Straw Concentrate	9670	Phenobarbital Pheniermine	2200
	Thebaine	9333	Prazepam	2764
	NON-NARCOTIC BASIC CLASSES	CODE	Quazépam Temazépam	2881 2925
	Amobarbital	2125	Triazolam	2007
	Amphetamine	1100	Zolpidem	2763
	Mothamphotamino Mothamphotatio	1105 1724		
	Methylphenidate Pentobarbital	2270	SCHEDULE V	
	Phencyclidine (PCP)	7471	OUNED ULE V	
	Phenmetrazine	1631		CODE
	Phenylacetone Secobarbital	6501 2315	Ordeles Oracle Descention (200esel400-1	0400
		2010	Codeline Cough Preparation (200mg/100ml or 100g)	9100
Notice in Registratio Mal	king Premant by Chack			

Notice to Registrants Making Payment by Check Authorization to Convert Your Check: If you send us a check to make your payment, your check will be converted into an electronic fund transfer. "Electronic fund transfer" is the term used to refer to the process in which we electronically instruct your financial institution to transfer funds from your account, rather than processing your check. By sending your completed, spind check to us, you authorize us to copy your check and to use the account information from your check and make an electronic fund transfer from your account for the same amount as the check. If the electronic fund transfer cannot be processed for technical reasons, you authorize us to process the copy of

your check. there are sufficient funds available in your checking account when you send us your check. If the electronic funds transfer cannot be completed because of insufficient funds, we may try to make the transfer up to two times.

Transaction between the dataset op to we then. Transaction hibromation: The electronic fund transfer from your account will be on the account statement you receive from your financial institution. However, the transfer may be in a different place on your statement than the place where your checks normally appear. For example, it may appear under "other withdrawals" or "other transactions." You will not receive your original check back from your financial institution. For security reasons, we will destroy your original check, but we will keep a copy of the check for record-keeping purposes. Your Abject: You should contact your financial institution immediately if you believe that the electronic fund transfer reported on your account statement was not properly authoritzed or is otherwise incorrect. Consumers have protections under Federal law called the Electronic Fund Transfer Act for an unauthorized or incorrect electronic fund transfer.

transfer.

		B NO 1117-0 DEA-224a (1-
INSTRUCTIONS	To renew by mail complete this application. Keep a copy for your records. Print clearly, using black or blue ink, or use a typewriter. Section 5 should be completed only if your information has changed. Mail this form to the address provided in Section 6 or use enclosed envelope. Include the correct payment amount. FEE IS NON-REFUNDABLE. If you have any questions call 800-882-9539 prior to submitting your application. Save time - renew online at www.deadiversion.usdoj.gov. IMPORTANT: DO NOT SEND THIS APPLICATION AND RENEW ONLINE.	ON :
	FEE IS NON-REFUNDABLE	
SECTION 1 DRUG SCHEDULES Check all that apply	Schedule II Narcotic Schedule III Narcotic Schedule IV Schedule II Non-Narcotic Schedule III Non-Narcotic Schedule V	
SECTION 2	Check this box if you need official order forms - for the purchase of schedule II narcotic/schedule II non-narcotic con	trolled substance
STATE LICENSE(S) Be sure to include both state license numbers if applicable	NO State License Number State Controlled Subs	
LIABILITY IMPORTANT: If you answered yes to thes question(5) on previous application, you must continue to answer yes and provide a statement of explanation. All questions in this section must be answered.	 B. Has the applicant ever been convicted of a crime in connection with controlled substances under state or federal law? C. Has the applicant ever surrendered (for cause) or had a federal controlled substance registration revoked, suspended, restricted, or denied? D. Has the applicant ever surrendered (for cause) or had a state professional license or controlled substance registration revoked, suspended, denied, restricted, or placed on probation? Is any such action pending? E. If the applicant is a corporation (other than a corporation whose stock is owned and traded by the public), association, partnership, or pharmacy, has any officer, partner, stockholder, or proprietor been convicted of a crime in connection with controlled substance sunder state or federal aw, or ever surrendered, for cause, or had a federal controlled substance registration revoked, suspended, denied, restricted or placed on probation? 	YES NO
SECTION 4		
EXPLANATION OF "YES" ANSWERS Applicants who have answered "YES" to questions B, C, D, or E above must provide a statement to explain such answers Use this space or attach	Date(s) of incident: Location(s) of incident: Nature of incident:	
a separate sheet and return with application	Result of incident:	

SECTION 5	Last Name (if registration is for individual) -OR- BUSINESS Name (if registration is for business)	
CHANGES TO APPLICANT		
IDENTIFICATION	First Name and Middle Initial	
DEBT COLLECTION		
INFORMATION	Tax Identification Number (if registration is for business) Social Security Number (if registration is for	individual)
Mandatory pursuant to Debt Collection Improvements Act		Provide SSN or TIN. See note #3 on bottom of page 2
	Address Line 1 (street address)	
IMPORTANT	Address Line 2	
Leave this section blank unless the		
registration information on	City State	e Zip Code
front page is incorrect.		
	B isiness Phone Number Business Fax Number	
SECTION 6	Make check payable to: Drug Enforcement Administration Check See page 4 of instructions for important information.	
METHOD OF PAYMENT	Check See page 4 of instructions for important information.	Mail this form with payment to:
Check one form of	American Express Discover Discover Visa	
payment only	Credit Card Number Expiration Date	U.S. Department of Justice Drug Enforcement Administration
		P.O. Box 105616
		Atlanta, GA 30348-5616
Sign if paying by credit card	Signature of Card Holder	FEE IS NON-REFUNDABLE
	Printed Name of Card Holder	
SECTION 7	Check this box if the applicant is a federal, state, or local government operated hospital, ins	
CERTIFICATION OF EXEMPTION	Be sure to enter the name and address of the exempt institution on address lines 1 and 2 in Sec current registration certificate.	tion 5, if it is not already on your
from application fee	The undersigned hereby certifies that the applicant named hereon is a federal, state or local government op and is exempt from payment of the application fee.	erated hospital, institution or official,
Provide the name and phone number of the certifying official	Signature of certifying official (other than applicant) Dat	e
	Print or type name and title of certifying official Tele	phone No. (required for verification)
SECTION 8	I certify that the foregoing information furnished on this application is true and correct.	
APPLICANT'S SIGNATURE		
Sign in ink	Signature of applicant	Date
	Print or type name and title of applicant	
	WARNING: Section 843(a)(4)(A) of Title 21, United States Code states that any person who knowingly or in fraudulent information in the application is subject to imprisonment for not more than four years, a fine of no	
1. No registration will be i	issued unless a completed application form has been received (21 CFR 1301.13). Paperwork Reduction Act of 1995, no person is required to respond to a collection of information unless it displ	ave a valid OMB control sumber. The
valid OMB control num the time for reviewing i	r aperwork reduction Add of 1990, no person is required to respond to a collection or ninformation unless it dispu- ber for this collection is 1117-0014. Public reporting burden for this collection of information is estimated to aver instructions, searching existing data sources, gathering and maintaining the data needed, and completing and re provements Act of 1990 (PL 104-134) requires that you furnish your Taxpayer Identifying Number and/or Social	age 12 minutes per response, including viewing the collection of information.
4. PRIVACY ACT INFOR	d for debt collection procedures should your fee become uncollectable. MATION	
AUTHORITY:	Section 302 and 303 of the Controlled Substances Act of 1970 (PL 91-513) and Debt Collection Impro taxpayer identifying number and/or social security number).	
PURPOSE: ROUTINE US	To obtain information required to register applicants pursuant to the Controlled Substances Act of 1970 EES: The Controlled Substances Act Registration Records produces special reports as required for statistic	
	information from this system are made to the following categories of users for the purposes stated: A. Other federal law enforcement and regulatory agencies for law enforcement and regulatory purpose B. State and local law enforcement and regulatory agencies for law enforcement and regulatory purpo	
EFFECT:	c. State and local law enforcement and regulatory agencies to hav enforcement and regulatory purpo C. Persons registered under the Controlled Substances Act (PL 91-513) for the purpose of verifying th Failure to complete form will preclude processing of the application.	e registration of customers.
	RENEWAL - Page 2	
	DENEMOE - Lage 2	

Form-224a		ION FOR REN ry Instructions and						
ADDITIONAL INSTRUCTIONS	SECTION 1. DRUG SCHEDULES - Applicants should check all drug schedules to be handled. However, applicants must still comply with state requirements; federal registration does not overrule state restrictions. Check the order form box only if you intend to purchase or to transfer schedule II controlled substances.							
	SECTION 2. ORDER FORMS - Order forms will be mailed to the registered address following issuance of a Certificate of Registration.							
	SECTION 3. STATE LICENSE(S) - Federal registration by DEA is based upon the applicant 's compliance with applicable state and local laws. Applicants should contact the local state licensing authority prior to completing this application. If your state requires a separate controlled substance number, provide that number on this application. If a state license has not yet been issued, indicate "Pending". If state licensing authority is not required, indicate "No".							
		SECTION 4. LIABILITY - Applicants must answer all four questions for the application to be accepted for processing. If you answered "Yes" to any question, provide an explanation in the space provided. If additional space is required, you may attach a separate sheet of paper.						
	reduc or nev is req numb	e data entry errors. Enter w phone numbers. Fee e uired; after the street add er (SSN) on record is con	- Entry of missing data or r changes in previously pro exempt individuals should I dress a post office box may rect. If renewing a busines s mandatory pursuant to	wided registration inform ist the name and addres be included. Individua sentity, a valid tax iden	mation, such as name ch ss of the fee exempt insti Is renewing should ensu htification number (TIN) n	ange, address correction, itution. A physical address re that the social security nust be supplied.		
	SECTION 6. METH Third	OD OF PAYMENT - India	ate the desired method of rawn on foreign banks will	payment. Make check not be accepted. FEE	s payable to "Drug Enfor S ARE NON-REFUNDAE	cement Administration". BLE.		
	SECTION 7. CERT opera	IFICATE OF EXEMPTION ted hospitals, institutions	N - Exemption from payme and officials. The applicar umber of the certifying offic	ent of application fee is I nt's superior or agency (imited to federal, state or officer must certify exemp	r local government		
			Aust be the original signatu					
CONTACT INFORMATION	1. INTERNET: 2. TELEPHONE: 3. WRITTEN INQUI	Headquarters RIES: Drug Enforcen P.O. Box 2808 Washington, D	n be found on our web site Call Center: (800) 882-953 ment Administration 33 0.C. 20038-8083 ww (800, 877, and 888 are	39	isdoj.gov			
	ATLANTA DIVISIO ATTN: Registration 75 Spring Street, S' Atlanta, GA 30303	N OFFICE	DETROIT DIVISION 431 Howard Street Detroit, MI 48226		PHILADELPHIA D William J. Green F 600 Arch Street, R Philadelphia, PA 1	ederal Building oom 10224		
	Georgia North Carolina	(888) 869-9935 (888) 219-8689	Kentucky Michigan Ohio	(800) 230-6844 (800) 230-6844 (800) 230-6844	Delaware Pennsylvania	(888) 393-8231 (888) 393-8231		
	South Carolina Tennessee BOSTON DIVISION JFK Federal Buildin	(888) 219-7898 SION OFFICE	EL PASO DIVISION El Paso Federal Just 600 South Mesa Hills El Paso, TX 79912	ice Center	PHOENIX DIVISIO 3010 N. 2nd Stree Phoenix, AZ 85012	t, Suite 301		
	15 New Sudbury St Boston, MA 02203-	bury Street, Room E400	New Mexico	(915) 832-6014	Arizona	(800) 741-0902		
	Connecticut Maine Massachusetts	(617) 557-2200 (888) 272-5174 (617) 557-2468	HOUSTON DIVISION 1433 West Loop Sou Houston, TX 77027-8	th, Suite 600	SAN DIEGO DIVIS 4560 Viewridge Av San Diego, CA 92	renue 123-1637		
	New Hampshire Rhode Island Vermont	(888) 272-5174 (617) 557-2200 (888) 272-5174	88) 272-5174 17) 557-2200 Texas (S. & Central) (800) 743-0595	California (Southern) (800) 284-1152 SAN FRANCISCO DIVISION OFFICE				
	CARIBBEAN DIVIS P.O. Box 2167 San Juan, PR 0092	ION OFFICE	LOS ANGELES DIV 255 East Temple Stre Los Angeles, CA 900	eet, 20th Floor		Avenue, 14th Floor		
	Puerto Rico	(787) 775-1766	California (S. Central Hawaii	(888) 415-9822	-	n) (888) 304-3251		
	U.S. Virgin Islands CHICAGO DIVISIO	(787) 775-1786 N OFFICE	Nevada Trust Territory	(888) 415-9822 (213) 894-2216	SEATTLE DIVISIO 400 Second Avenu Seattle, WA 98119	ue, West		
	Kluczynski Federal 230 S. Dearborn St Chicago, IL 60604	Building	MIAMI DIVISION OF 8400 N.W. 53rd Stree Miami, FL 33166	FICE et	Alaska Idaho Oregon	(888) 219-4261 (888) 219-4261 (888) 219-4261		
	Illinois Indiana	(312) 353-1234 (312) 353-1236	Florida	(305) 590-4880	Washington	(888) 219-1418		
	Minnesota North Dakota Wisconsin	(312) 353-9166 (312) 353-9166 (312) 353-1236	NEWARK DIVISION 80 Mulberry Street, 2 Newark, NJ 07102		ST. LOUIS DIVISIO 317 South 16th Str St. Louis, MO 6310	reet		
	DALLAS DIVISION 10160 Technology & Dallas, TX 75220		New Jersey NEW ORLEANS DIV 3838 N. Causeway B	(888) 356-1071 (ISION OFFICE	lowa Kansas Missouri Nebraska	(888) 803-1179 (888) 803-1179 (888) 803-1179 (888) 803-1179 (888) 803-1179		
	Oklahoma Texas (Northern)	(888) 336-4704 (888) 336-4704	Lakeway III, Suite 18 Metairie, LA 70002		South Dakota	(888) 803-1179		
	DENVER DIVISION 115 Inverness Drive Englewood, CO 80	e, East	Alabama Arkansas Louisiana Mississiani	(888) 514-8051 (888) 514-7302 (888) 514-7302 (888) 514-7302	WASHINGTON, D Techworld Plaza 800 K Street, N.W. Washington, D.C.			
	Colorado Montana Utah Wyoming	(800) 328-6900 (800) 326-6900 (800) 326-6900 (800) 326-6900	Mississippi NEW YORK DIVISIO 99 Tenth Avenue New York, NY 10011	(888) 514-7302 N OFFICE	District of Columbi Maryland Virginia West Virginia	a (877) 801-7974 (877) 330-6670 (877) 801-7974 (877) 330-6670		
			New York	(877) 883-5789 (212) 337-1593 (212) 337-1594				
RENEWAL INST - Page 3	1							

SCHEDULE I		SCHEDULE III	
NARCOTIC & NON-NARCOTIC BASIC CLASSES	CODE	NARCOTIC BASIC CLASSES	COD
	9319	Buprenorphine	9064 9319
Acetorphine Acetylmethadol	9601	Codeine up to 90 mg/du plus other ingredients Dihydrocodeineup to 90 mg/du plus other ingredients	9807
Allylprodine	9602	Ethylmorphine up to 15 mg/du plus other ingredients	9808
Alphacetylmethadol (except LAAM)	9603	Hydrocodone up to 15 mg/du plus other ingredients Marshing up to 50 mg/100ml or am plus other ingred	9806 9810
Bufotenine Dextromoramide	7433 9613	Morphine up to 50 mg/100ml or gm plus other ingred. Opium up to 500 mg/100m. plus other active ingred.	9809
Diethyltryptamine (DET)	7434		
2,5 - Dimethoxyamphetamine (DMA)	7396	NON-NARCOTIC BASIC CLASSES	COL
Dimethyltryptamine (DMT)	7435	Apphalia Staroida	4000
Etorphine (except hydrochloride salt) gamma-Hydroxybutyric acid (except drug product)	9056 2010	Anabolic Steroids Benzphetamine	122
Heroin	9200	Butalbital	2100
Ibogaine	7260	Dronabinol Pharmaceutical Product	736
Ketobemidone	9628	GHB Drug Product (gamma-Hydroxybutyric acid)	2010
Lysergic acid diethylamide (LSD) Marihuana	7315 7360	Ketamine Methyprylon	728 257
Mannuaria Mescaline	7381	Pentobarbital plus noncontrolled active ingredients	227
Methagualone	2565	Pentobarbital suppository	227
3,4 - Methylenedioxyamphetamine (MDA)	7400	Phendimetrazine	161
3.4 - Methylenedioxymethamphetamine (MDMA)	7405	Secobarbital plus noncontrolled active ingredients	231
n- Ethyl - 1 - Phenylcyclohexylamine (PCE) Peyote	7455 7415	Secobarbital suppository Thiopental	231 232
1 - (1-Phenylcyclohexyl)pyrrolidine (PCP)	7458	Vinbarbital	233
Psilocybin	7437		200
Psilocyn	7438		
Tetrahydrocannabinols (THC) 1-[1-(2-Thienyl)-cyclohexyl]-piperidine	7370 7470	SCHEDULE IV	
		NARCOTIC BASIC CLASSES	COL
SCHEDULE II		Dextropropoxyphene du Difenoxin 1mg/25ug atropine SO4/du	9278 9163
NARCOTIC BASIC CLASSES	CODE	NON-NARCOTIC BASIC CLASSES	COL
Alphaprodine	9010	Alexandress	288
Anileridine	9020	Alprzolam Barbital	200.
Cocaine Codeine	9041 9050	Chloral Hydrate	246
Dextropropoxyphene (bulk)	9273	Chlordiazepoxide	274
Diphenoxylate	9170	Clorazepate	276
Diprenorphine (M50-50)	9058	Diazepam Diethylpropion	276 161
Ethylmorphine	9190 9059	Fenfluramine	167
Etorphine Hydrochloride (M-99) Glutethimide	2550	Flurazepam	276
Hydrocodone	9193	Halazepam	276
Hydromorphone	9150	Lorazepam	288
Levo-alphacetylmethadol (LAAM)	9648	Mazindol Mebutamate	160 280
Levorphanol Meperidine	9220 9230	Mephobarbital (Methylphenobarbital)	225
Methadone	9250	Meprobamate	282
Morphine	9300	Methohexital	226
Opium, powdered	9639	Midazolam	288
Opium, raw	9600	Oxazepam Paraldehyde	283 258
Oxycodone Oxymorphone	9143 9652	Pemoline	1530
Poppy Straw	9671	Pentazocine	970
Poppy Straw Concentrate	9670	Phenobarbital	228
Thebaine	9333	Phentermine	1640 2764
NON NADOOTIO BASIO OLASSES	0005	Prazepam Quazepam	288
NON-NARCOTIC BASIC CLASSES	CODE	Temazepam	292
Amobarbital	2125	Triazolam	2887
Amphetamine	1100	Zolpidem	278
Methamphetamine	1105		
Methylphenidate Bostobarbital	1724		
Pentobarbital Phencyclidine (PCP)	2270 7471	SCHEDULE V	
Phenmetrazine	1631		COL
Phenylacetone	8501		001
Secobarbital	2315	Codeine Cough Preparation (200mg/100ml or 100g)	9100

the term used to relet to the process in which we electronically instruct your infancial instruction to finance funds from your account to our account, rather than processing your oheck. By sending your completed, signed check to us, you authorize us to copy your check and to use the account information from your check to make an electronic fund transfer from your account for the same amount as the check. If the electronic fund transfer cannot be processed for technical reasons, you authorize us to process the copy of your check.

Insufficient Funds: The electronic funds transfer from your account will usually occur with 24 hours, which is faster than a check is normally processed. Therefore, make sure

there are sufficient funds available in your checking account when you send us your check. If the electronic funds transfer cannot be completed because of insufficient funds, we may try to make the transfer up to two times. Transaction Information: The electronic fund transfer from your account will be on the account statement you receive from your financial institution. However, the transfer may be in a different place on your statement than the place where your checks normally appear. For example, it may appear under "other withdrawals" or "other transactions." You will not receive your original check back from your financial institution. For security reasons, we will destroy your original check, but we will keep a copy of the check for record-keeping purposes.

Your Rights: You should contact your financial institution immediately if you believe that the electronic fund transfer reported on your account statement was not properly authorized or is otherwise incorrect. Consumers have protections under Federal law called the Electronic Fund Transfer Act for an unauthorized or incorrect electronic fund transfer.

RENEW AL INST - Page 4

Form-363	APPLICATION FOR REGISTRATION Under the Narcotic Addict Treatment Act of 1974	APPROVED OMB NO 1117-0015 FORM DEA-363 (11-05) Previous editions are obsolete
INSTRUCTIONS	 To apply by mail complete this application. Keep a copy for your records. Frint clearly using black or blue ink, or use a typewriter. Section 1 should be completed only if your information has changed. Mail this form to the address provided in Saction 8 or use enclosed envelope. Include the correct payment amount. FEE IS NON-REFUNCABLE. If you have any questions contact 800-882-953 prior to sumting your application. Save time - apply online at www.deadiversion.usdoj.gov. 	REGISTRATION INFORMATION :
	IMPORTANT: DO NOT SEND THIS APPLICATION AND APPLY ONLINE.	
		Fee for 1 year is \$130 FEE IS NON-REFUNDABLE
SECTION 1 APPL	ICANT	
	TIFICATION	
Business or Facility	Name (If registration is for business entity or is fee exempt)	
Business or Facility	Name 2 ("doing business as", continuation of business name, or name of fee e	xempt institution)
Address Line 1 (stre	eet address)	
Address Line 2		
City		State Zip Code
Business Phone Nu	mber Business Fax Number	
DEBT COLLECTION	Tax Identification Number	
Mandatory pursuant to Debt Collection Improvements Act		See note #3 on bottom of page 2.
SECTION 2	NTP - Maintenance NTP - Co	mpounder / Maintenance
BUSINESS ACTIVITY Check one box only	NTP - Detoxification	mpounder / Detoxification
	NTP - Maintenance and Detoxification	mpounder / Maintenance and Detoxification
SECTION 3	Schedule II Schedul	e III
DRUG SCHEDULES	Check this box if you require official order forms - for purchase or transfer of	schedule II controlled substances
Check all that apply		
	re you currently authorized by the Food and Drug Administration for the busines	ss activity described in this application?
FDA PERMIT Mandatory for approval	YES PENDING NO	FDA Number
SECTION 5 Are the s	you currently authorized to prescribe, distribute, dispense, conduct research, or schedules for which you are applying under the laws of the state or jurisdiction i	otherwise handle the controlled substances in n which you are operating or propose to operate?
STATE LICENSE(S)	YE8, I have a licence	State
	NOT REQUIRED by this state	License Number
	NEW - Page 1	

SECTION 6	1. Has the applicant ever been convicted of a crime in connection with controlled substances una	ier state or federal law?	YES	
LIABILITY	Has the applicant ever surrendered (for cause) or had a federal controlled substance registration restricted, or denied?	n revoked, suspended,		
IMPORTANT: All questions in	3. Has the applicant ever surrendered (for cause) or had a state professional license or controlled revoked, suspended, denied, restricted, or placed on probation? Is any such action pending?	substance registration		
this section must be answered.	4. If the applicant is a corporation (other than a corporation whose stock is owned and traded by partnership, or pharmacy, has any officer, partner, stockholder, or proprietor been convicted of controlled substances under statie or federal law, or ever surrenderined, for cause, or had a feder registration revoked, suspended, restricted, denied, or ever had a state professional license or or registration revoked, suspended, denied, restricted or placed on probation?	the public), association, a crime in connection with al controlled substance		
EXPLANATION OF "YES" ANSWERS	Date(s) of Incident: Location(s) of Incident:			
Applicants who have answered "YES" to any of the four question above must provide a statement to explain such answers				
Use this space or attact a separate sheet and return with application	h Result of incident:			
SECTION 7 CERTIFICATION OF EXEMPTION from application fee	Check this box if the applicant is a federal, state, or local government-operated narcotic treat Be sure to enter name and address of the exempt institution in Section 1. The undersigned hereby certifies that the applicant named hereon is a federal, state or local government program, and is exempt from payment of the application fee.		2	
Provide the name and phone number of the certifying official	Signature of certifying official (other than applicant) Date			
cardiying cincar	Print or type name and title of certifying official Telep	hone No. (required for vertile	ation)	
SECTION 8 METHOD OF PAYMENT Check one form of payment only	Check Make Check payable to: Drug Enforcement Administration See page 3 of instructions for Important Information. American Express Discover Master Card Visa	Mail this form with p U.S. Department o		
	Credit Card Number Expiration Date	Drug Enforcement Ad P.O. Box 280 Washington DC 20	ministr 63	ation
Sign If paying by credit card	Signature of Card Holder	FEE IS NON-REFU	NDAB	LE
	Printed Name of Card Holder			
SECTION 9	I certify that the foregoing information furnished on this application is true and correct.			
SIGNATURE Sign in Ink	Signature of applicant D	ate		
	Print or type name and title of applicant WARNING: Section 643(a)(4)(A) of Title 21, United States Code states that any person who knowingly or inte fraudulent information in the application is subject to imprisonment for not more than four years, a fine of notin			
3. The Debt Collection		eviewing the collection of infor Security Number on this appli	mation. cation.	
PURPOSE: ROUTINE U	texpayer identifying number and/or social security number). To obtain information required to register applicants pursuant to the Controlled Substances Act of 197	0. ai analytical purposes. Disclo es. oses.		
EFFECT:	Failure to complete form will preclude processing of the application. NEW - Page 2			

Form-363	APPLICATION FOR REGISTRATION Supplementary Instructions and Information
ADDITIONAL INSTRUCTIONS	SECTION 1. APPLICANT IDENTIFICATION - Information must be typed or printed in the blocks provided to help reduce data entry errors.
	Fee exempt applicant should list the name and address of the fee exempt institution. A physical address is required; a post office box may be included after the street address.
	Applicant must enter a valid tax identification number (TIN). Debt collection information is mandatory pursuant to the Debt Collection Improvement Act of 1996.
	SECTION 2. BUSINESS ACTIVITY. Indicate only one.
	SECTION 3. DRUG SCHEDULES - Applicant should check all drug schedules to be handled. However, applicant must still comply with state requirements; federal registration does not overrule state restrictions.
	Check the order form box only if you intend to purchase or to transfer schedule II controlled substances. Order forms will be mailed to the registered address following issuance of a Certificate of Registration.
	SECTION 4. FDA PERMIT - Authorization by the Food & Drug Administration is mandatory for DEA Registration approval. Enter the status of your FDA authorization and the FDA number.
	SECTION 5. STATE LICENSE(S) - Federal registration by DEA is based upon the applicant 's compliance with applicable state and local laws.
	Applicant should contact the local state licensing authority prior to completing this application. Check that you are currently authorized by the state and provide your state license number. If state licensing is not required, indicate "Not required by this state".
	SECTION 6. LIABILITY - Applicant must answer all four questions for the application to be accepted for process
	If you answered "Yes" to any question, provide an explanation in the space provided. If additional space is required, you may attach a separate sheet of paper.
	SECTION 7. CERTIFICATE OF EXEMPTION - Exemption from payment of application fee is limited to federal, state or local government-operated narcotic treatment program.
	The applicant's superior or agency officer must certify exempt status. The signature, authority title, and telephone number of the certifying official (other than the applicant) must be provided.
	SECTION 8. METHOD OF PAYMENT - Indicate the desired method of payment. Make checks payable to "Drug Enforcement Administration". Third-party checks or checks drawn on foreign banks will not be accepted.
	FEES ARE NON-REFUNDABLE.
	SECTION 9. APPLICANT'S SIGNATURE - Must be the original signature (in ink) of the applicant.

Notice to Registrants Making Payment by Check

Authorization to Convert Your Check: If you send us a check to make your payment, your check will be converted into an electronic fund transfer. "Electronic fund transfer" is the term used to refer to the process in which we electronically instruct your financial institution to transfer funds from your account to our account, rather than processing your check. By sending your completed, signed check to us, you authorize us to copy your check and to use the account information from your check to make an electronic fund transfer from your account for the same amount as the check. If the electronic fund transfer cannot be processed for technical reasons, you authorize us to process the copy of your check.

Insufficient Funds: The electronic funds transfer from your account will usually occur with 24 hours, which is faster than a check is normally processed. Therefore, make sure there are sufficient funds available in your checking account when you send us your check. If the electronic funds transfer cannot be completed because of insufficient funds, we may try to make the transfer up to two times.

Transaction Information: The electronic fund transfer from your account will be on the account statement you receive from your financial institution. However, the transfer may be in a different place on your statement than the place where your checks normally appear. For example, it may appear under "other withdrawals" or "other transactions." You will not receive your original check back from your financial institution. For security reasons, we will destroy your original check, but we will keep a copy of the check for record-keeping purposes.

Your Rights: You should contact your financial institution immediately if you believe that the electronic fund transfer reported on your account statement was not properly authorized or is otherwise incorrect. Consumers have protections under Federal law called the Electronic Fund Transfer Act for an unauthorized or incorrect electronic fund transfer.

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orm-363		Instructions and Information				
CONTACT	1. INTERNET:	Information	can be found on our web	site at www.deadiversion.	usdoj.gov	
NFORMATION	2. TELEPHONE:	Headquarte	ers Call Center: (800) 882-	9539		
	3. WRITTEN INQUI	RIES: Drug Enfor	cement Administration			
		P.O. Box 2 Washingto	cement Administration 8083 n DC 20038-8083			
	4. DEA OFFICES: D	EA Offices are listed l	below (800, 877, and 888 a	are toll-free numbers).		
ATLANTA DIVISI		DETROIT DIVISIO		PHILADELPHIA DIV		
ATTN: Registratio 75 Spring Street,		431 Howard Stree Detroit, MI 48226		William J. Green Fed 600 Arch Street, Roo		
Atlanta, GA 30303				Philadelphia, PA 191		
Ceorgia	(888) 869-9935	Kentucky	(800) 230-6844 (800) 230-6844	Delaware	(888) 393-8231	
Georgia North Carolina	(888) 219-8689	Michigan Ohio	(800) 230-6844	Pennsylvania	(888) 393-8231	
South Carolina	(866) 533-6983	EL PASO DIVISIO	NOFFICE		OFFICE	
Tennessee	(888) 219-7898	EL PASO DIVISIO El Paso Federal J		PHOENIX DIVISION 3010 N. 2nd Street, 5		
BOSTON DIVISIO		600 South Mesa H	-lills Drive, Suite 2000	Phoenix, AZ 85012		
JFK Federal Build	ing Street, Room E400	El Paso, TX 7991	2	Arizona	(800) 741-0902	
Boston, MA 02203	3-0131	New Mexico	(915) 832-6014		(,	
Connections	(047) 557 0000	HOUETON DRAG	. ,	SAN DIEGO DIVISIO		
Connecticut Maine	(617) 557-2200 (888) 272-5174	HOUSTON DIVIS 1433 West Loop S		4560 Viewridge Avenue San Diego, CA 92123-1637		
Massachusetts	(617) 557-2468	Houston, TX 7702	27-9506			
New Hampshire Rhode Island	(888) 272-5174 (617) 557-2200	Texas (S. & Centr	al) (800) 743-0595	California (Southern	(800) 284-1152	
Vermont	(888) 272-5174			SAN FRANCISCO DIVISI		
	. ,	LOS ANGELES DIVISION OFFICE 450 Golden Gate Avenue, 14th			enue, 14th Floor	
CARIBBEAN DIVISION OFFICE P.O. Box 2167		Los Angeles, CA	255 East Temple Street, 20th Floor Los Angeles, CA 90012		P.O. Box 36035 San Francisco, CA 94102	
San Juan, PR 009	22-2167					
Puerto Rico	(787) 775-1766	California (S. Cen Hawaii	tral) (213) 621-6960 (888) 415-9822	California (Northern)	(888) 304-3251	
U.S. Virgin Islands		Nevada (888) 415-9822		SEATTLE DIVISION		
	ON OFFICE	Trust Territory	(213) 894-2216	400 Second Avenue, Secttle, WA 09110	West	
CHICAGO DIVISI Kluczynski Federa	al Building	MIAMI DIVISION	OFFICE	Seattle, WA 98119		
230 S. Dearborn S	Street, Suite 1200	8400 N.W. 53rd S		Alaska	(888) 219-4261	
Chicago, IL 60604	L	Miami, FL 33166		ldaho Oregon	(888) 219-4261 (888) 219-4261	
Illinois	(312) 353-1234	Florida	(305) 590-4880	Washington	(888) 219-1418	
Indiana Minnesota	(312) 353-1236 (312) 353-9166	NEWARK DIVISI	ON OFFICE	ST. LOUIS DIVISION	OFFICE	
Minnesota North Dakota	(312) 353-9166 (312) 353-9166	80 Mulberry Stree		317 South 16th Stree		
Wisconsin	(312) 353-1236	Newark, NJ 07102	2	St. Louis, MO 63103		
DALLAS DIVISIO 10160 Technology		New Jersey	(888) 356-1071	lowa Kansas	(888) 803-1179 (888) 803-1179	
Dallas, TX 75220	Endig Edot		DIVISION OFFICE	Missouri	(888) 803-1179	
Oklahoma	(888) 336-4704	3838 N. Causewa Lakeway III, Suite	y Blvd	Nebraska South Dakota	(888) 803-1179 (888) 803-1179	
Texas (Northern)		Metairie, LA 7000			1 7	
DENVER DIVISIO	NOFFICE	Alabama	(888) 514-8051	WASHINGTON, D.C. Techworld Plaza	DIVISION OFFIC	
115 Inverness Driv	ve, East	Arkansas	(888) 514-7302	800 K Street, N.W., S		
Englewood, CO 8		Louisiana	(888) 514-7302	Washington, D.C. 20		
Colorado	(800) 326-6900	Mississippi	(888) 514-7302	District of Columbia	(877) 801-7974	
Montana	(800) 326-6900		NEW YORK DIVISION OFFICE		(877) 330-6670	
Utah Wyoming	(800) 326-6900 (800) 326-6900	99 Tenth Avenue New York, NY 100	011	Virginia West Virginia	(877) 801-7974 (877) 330-6670	
myoning	(500) 520-6800			meac virginita	(011) 330-0010	
		New York	(877) 883-5789 (212) 337-1593			
			(212) 337-1593			

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Form-363a	RENEWAL APPLICATION FOR REGISTRATION Under the Narcotic Addict Treatment Act of 1974	APPROVED OMB NO 1117-0015 FORM DEA-363a (11-05) Previous editions are obsolete
INSTRUCTIONS	 To apply by mail complete this application. Keep a copy for your records. Print clearly, using black or blue ink, or use a typewriter. Section 1 should be completed only if your information has changed. Mail this form to the address provided in Section 7 or use enclosed envelope. Include the correct payment amount. FEE IS NON-REFUNDABLE. If you have any questions contact 800-882-9639 prior to submitting your application. Save time - renew online at www.deadiversion.usdoj.gov. 	REGISTRATION INFORMATION : DEA # REGISTRATION EXPIRES
	IMPORTANT: DO NOT SEND THIS APPLICATION AND APPLY ONLINE.	
		FEE IS NON-REFUNDABLE
	ICANT TIFICATION	
Business or Facility	Name (if registration is for business entity or is fee exempt)	
Business or Facility	Name 2 ("doing business as", continuation of business name, or name of fee ex	xempt institution)
Address Line 1 (stre	eet address)	
Address Line 2		
City		State Zip Code
Business Phone Nu	mb. r Bu: ness F ax Number	
DEBT COLLECT ON		
INFORMATION	Tax Identification Number	
Mandatory pursuant to Debt Collection Improvements Act		See note #3 on bottom of page 2.
SECTION 2	Schedule II Schedule III	
DRUG SCHEDULES		
Check all that apply	Check this box if you require official order forms - for purchase or transfer of so	hedule II controlled substances.
SECTION 3 A	re you currently authorized by the Food and Drug Administration for the busines	activity described in this application?
FDA PERMIT YI Mandatory for approval	ES PENDING NO	FDA Number
Mandatory for approval		T DA Number
SECTION 4 Are the s	you currently authorized to prescribe, distribute, dispense, conduct research, or chedules for which you are applying under the laws of the state or jurisdiction in	otherwise handle the controlled substances in n which you are operating or propose to operate?
STATE LICENSE(S)	YES, I have a license	State License Number
1	NOT REQUIRED by this state	
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		YES NO
	1. Has the applicant ever been convicted of a crime in connection with controlled substances	
	2. Has the applicant ever surrendered (for cause) or had a federal controlled substance registr restricted, or denied?	ration revoked, suspended,
IMPORTANT: All questions in this section must	3. Has the applicant ever surrendered (for cause) or had a state professional license or contro revoked, suspended, denied, restricted, or placed on probation? Is any such action pendi	lled substance registration
be answered.	4. If the applicant is a corporation (other than a corporation whose stock is owned and traded partnership, or pharmacy, has any officer, partner, stockholder, or proprietor been convicted controlled substances under state or federal law, or ever surrendered, for cause, or had a fe registration revoked, suspended, restricted, denied, or ever had a state professional license registration revoked, suspended, denied, réstricted or placed on probation?	of a crime in connection with
EXPLANATION OF "YES" ANSWERS	Date(s) of incident: Location(s) of incident:	
Applicants who have answered "YES" to any of the four questions above must provide a statement to explain such answers	Nature of incident:	
Use this space or attach a separate sheet and return with application	Result of incident:	
SECTION 6 CERTIFICATION OF EXEMPTION from application fee	Check this box if the applicant is a federal, state, or local government-operated narcotic f Be sure to enter name and address of the exempt institution in Section 1. The undersigned hereby certifies that the applicant named hereon is a federal, state or local treatment program, and is exempt from payment of the application fee.	
Provide the name and phone number of the certifying official	Signature of certifying official (other than applicant)	ate
	Print or type name and title of certifying official	elephone No. (required for verification)
SECTION 7 METHOD OF PAYMENT Check one form of	Check Make check payable to: Drug Enforcement Administration See page 3 of instructions for important information. American Express Discover Master Card Visa	Mail this form with payment to:
payment only	Credit Card Number Expiration Date	U.S. Department of Justice Drug Enforcement Administration P.O. Box 28083 Washington DC 20038-8083
Sign if paying by credit card	Signature of Card Holder	FEE IS NON-REFUNDABLE
	Printed Name of Card Holder	
SECTION 8	I certify that the foregoing information furnished on this application is true and correct.	
APPLICANT'S SIGNATURE Sign in ink	Signature of applicant	Date
5	Print or type name and title of applicant	
	WARNING: Section 843(a)(4)(A) of Title 21, United States Code states that any person who knowingly or fraudulent information in the application is subject to imprisonment for not more than four years, a fine of	intentionally furnishes false or not more than \$30,000, or both.
 In accordance with the valid OMB control nur the time for reviewing The Debt Collection Ir 	5: Section 302 and 303 of the Controlled Substances Act of 1970 (PL 91-513) and Debt Collection I taxpayer identifying number and/or social security number). To obtain information required to register applicants pursuant to the Controlled Substances Act of SES: The Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registration Records produces special reports as required for statement of the Controlled Substances Act Registratement of the Controlled Substances Act Registratement of the C	a verage 30 minutes per response, including ind reviewing the collection of information. ocial Security Number on this application. mprovements Act of 1998 (PL 104-134) (for 1970. tistical analytical purposes. Disclosures of
EFFECT:	information from this system are made to the following categories of users for the purposes state A. Other federal law enforcement and regulatory agencies for law enforcement and regulatory or B. State and local law enforcement and regulatory agencies for law enforcement and regulatory or C. Persons registered under the Controlled Substances Act (PL 91-513) for the purpose of verify Failure to complete form will preclude processing of the application. RENEWAL - Page 2	d: rposes. purposes.

Form-363a	APPLICATION FOR RENEWAL Supplementary Instructions and Information
ADDITIONAL INSTRUCTIONS	SECTION 1. APPLICANT IDENTIFICATION - Entry of missing data or corrections ONLY must be typed or printed in the blocks provided to help reduce data entry errors. Enter changes in previously provided registration information, such as name change, address correction, or new phone numbers.
	Fee exempt applicant should list the name and address of the fee exempt institution.
	A physical address is required; a post office box may be included after the street address.
	Applicant should ensure that the tax identification number (TIN) on record is correct. Debt collection information is mandatory pursuant to the Debt Collection Improvement Act of 1996.
	SECTION 2. DRUG SCHEDULES - Applicant should check all drug schedules to be handled. However, applicants must still comply with state requirements; federal registration does not overrule state restrictions.
	Check the order form box only if you intend to purchase or to transfer schedule II controlled substances. Order forms will be mailed to the registered address following issuance of a Certificate of Registration renewal.
	SECTION 3. FDA PERMIT - Authorization by the Food & Drug Administration is mandatory for DEA Registration approval. Enter the status of your FDA authorization and the FDA number.
	SECTION 4. STATE LICENSE(S) - Federal registration by DEA is based upon the applicant 's compliance with applicable state and local laws.
	Applicant should contact the local state licensing authority prior to completing this application. Check that you are currently authorized by the state and provide your state license number. If state licensing is not required, indicate "Not required by this state".
	SECTION 5. LIABILITY - Applicant must answer all four questions for the application to be accepted for processin
	If you answered "Yes" to any question, provide an explanation in the space provided. If additional space is required, you may attach a separate sheet of paper.
	SECTION 6. CERTIFICATE OF EXEMPTION - Exemption from payment of application fee is limited to federal, state or local government-operated narcotic treatment program.
	The applicant's superior or agency officer must certify exempt status. The signature, authority title, and telephone number of the certifying official (other than the applicant) must be provided.
	SECTION 7. METHOD OF PAYMENT - Indicate the desired method of payment. Make checks payable to "Drug Enforcement Administration". Third-party checks or checks drawn on foreign banks will not be accepted.
	FEES ARE NON-REFUNDABLE.
	SECTION 8. APPLICANT'S SIGNATURE - Must be the original signature (in ink) of the applicant.

Notice to Registrants Making Payment by Check

Authorization to Convert Your Check: If you send us a check to make your payment, your check will be converted into an electronic fund transfer. "Electronic fund transfer" is the term used to refer to the process in which we electronically instruct your financial institution to transfer funds from your account to our account, rather than processing your check. By sending your completed, signed check to us, you authorize us to copy your check and to use the account information from your check to make an electronic fund transfer from your account for the same amount as the check. If the electronic fund transfer cannot be processed for technical reasons, you authorize us to process the copy of your check.

Insufficient Funds: The electronic funds transfer from your account will usually occur with 24 hours, which is faster than a check is normally processed. Therefore, make sure there are sufficient funds available in your checking account when you send us your check. If the electronic funds transfer cannot be completed because of insufficient funds, we may try to make the transfer up to two times.

Transaction Information: The electronic fund transfer from your account will be on the account statement you receive from your financial institution. However, the transfer may be in a different place on your statement than the place where your checks normally appear. For example, it may appear under "other withdrawals" or "other transactions." You will not receive your original check back from your financial institution. For security reasons, we will destroy your original check, but we will keep a copy of the check for record-keeping purposes.

Your Rights: You should contact your financial institution immediately if you believe that the electronic fund transfer reported on your account statement was not properly authorized or is otherwise incorrect. Consumers have protections under Federal law called the Electronic Fund Transfer Act for an unauthorized or incorrect electronic fund transfer.

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Form-363a	APPLICATION Supplementary In					
CONTACT	1. INTERNET:	Infor	mation can be found on our we	b site at www.deadiversior	n.usdoj.gov	
NFORMATION	2. TELEPHONE:	Hea	dquarters Call Center: (800) 882	Call Center: (800) 882-9539		
	3. WRITTEN INQU	RIES: Druc P.O.	Enforcement Administration Box 28083 hington DC 20038-8083			
	4. DEA OFFICES: [listed below (800, 877, and 888	are toll-free numbers).		
ATLANTA DIVIS ATTN: Registrati 75 Spring Street Atlanta, GA 3030	on , SW, Suite 800	DETROIT I 431 Howar Detroit, MI		PHILADELPHIA DI William J. Green Fe 600 Arch Street, Ro Philadelphia, PA 19	deral Building om 10224	
Georgia North Carolina	(888) 869-9935 (888) 219-8689	Kentucky Michigan Ohio	(800) 230-6844 (800) 230-6844 (800) 230-6844	Delaware Pennsylvania	(888) 393-823 (888) 393-823	
South Carolina Tennessee BOSTON DIVISI JFK Federal Buil		El Paso Fe	DIVISION OFFICE deral Justice Center Mesa Hills Drive, Suite 2000 (79912	PHOENIX DIVISIOI 3010 N. 2nd Street, Phoenix, AZ 85012		
	Street, Room E400	New Mexic		Arizona	(800) 741-09	
Connecticut Maine	(617) 557-2200 (888) 272-5174	HOUSTON	o (915) 832-6014 DIVISION OFFICE Loop South, Suite 600	SAN DIEGO DIVIS 4560 Viewridge Ave San Diego, CA 921	enue	
Massachusetts New Hampshire	(617) 557-2468 (888) 272-5174	Houston, T	X 77027-9506	California (Southerr		
Rhode Island Vermont	(617) 557-2200 (888) 272-5174	Texas (S. 8	Central) (800) 743-0595	SAN FRANCISCO	. ,	
CARIBBEAN DIVISION OFFICE P.O. Box 2167		255 East Te	ELES DIVISION OFFICE emple Street, 20th Floor s, CA 90012	450 Golden Gate Av P.O. Box 36035	450 Golden Gate Avenue, 14th Floor	
San Juan, PR 00	California (S. Central) (213) 621-6960		California (Northern) (888) 304-32		
Puerto Rico (787) 775-1766 U.S. Virgin Islands (787) 775-1766		Hawaii Nevada Trust Territo	(888) 415-9822 (888) 415-9822 ory (213) 894-2216	SEATTLE DIVISION OFFICE 400 Second Avenue, West		
CHICAGO DIVIS Kluczynski Fede 230 S. Dearborn Chicago, IL 6060	ral Building Street, Suite 1200		ISION OFFICE 53rd Street 33166	Seattle, WA 98119 Alaska Idaho	(888) 219-426 (888) 219-426	
Illinois	(312) 353-1234	Florida	(305) 590-4880	Oregon Washington	(888) 219-426 (888) 219-141	
Indiana Minnesota North Dakota Wisconsin	(312) 353-1236 (312) 353-9166 (312) 353-9166 (312) 353-9166 (312) 353-1236	NEWARK I 80 Mulberry Newark, N	DIVISION OFFICE y Street, 2nd Floor J 07102	ST. LOUIS DIVISIO 317 South 16th Stre St. Louis, MO 6310	et	
DALLAS DIVISI		New Jersey	(888) 356-1071	lowa Kansas	(888) 803-1179 (888) 803-1179	
10160 Technolog Dallas, TX 75220	jy Divu., East)		EANS DIVISION OFFICE	Kansas Missouri Nebraska	(888) 803-1179	
Oklahoma Texas (Northern)	(888) 336-4704 (888) 336-4704	3838 N. Ca Lakeway III Metairie, L/	useway Blvd I, Suite 1800 A 70002	South Dakota	(888) 803-1179 (888) 803-1179	
DENVER DIVISI 115 Inverness Di Englewood, CO	rive, East	Alabama Arkansas Louisiana Mississippi	(888) 514-8051 (888) 514-7302 (888) 514-7302 (888) 514-7302 (888) 514-7302	WASHINGTON, D.0 Techworld Plaza 800 K Street, N.W., Washington, D.C. 2	Suite 500	
Colorado Montana Utah Wyoming	(800) 326-6900 (800) 326-6900 (800) 326-6900 (800) 326-6900 (800) 326-6900		K DIVISION OFFICE	District of Columbia Maryland Virginia West Virginia	(877) 801-797 (877) 330-667((877) 801-797 (877) 330-667(
		New York	(877) 883-5789 (212) 337-1593 (212) 337-1594			

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