A drug distribution system and method utilizes a central pharmacy and database to track all prescriptions for a sensitive drug. Information is kept in the database regarding all physicians allowed to prescribe the sensitive drug, and all patients receiving the drug. Abuses are identified by monitoring data in the database for prescription patterns by physicians and prescriptions obtained by patients. Further verification is made that the physician is eligible to prescribe the drug, by consulting a separate database, and optionally whether any actions are taken against the physician. Multiple controls beyond those for normal drugs are imposed on the distribution depending on the sensitivity of the drug.
Peripheral and Central Nervous System Drugs Advisory Committee*, Department of Health and Human Services Food and Drug Administration Center for Drug Evaluation and Research, Holiday Ina, Bethesda, Maryland,(Jun. 6, 2001),7 pages.


* cited by examiner
FIG. 1
MD MAILS/FAXES IN Rx/ENROLLMENT FORM

AN INTAKE/REIMBURSEMENT SPECIALIST MAKE A COPY OF THE Rx/ENROLLMENT FORM (THE COPY IS STAMPED "COPY") AND THE ORIGINAL FAX IS THEN FORWARDED TO THE PHARMACY TEAM

MD MAILS/FAXES IN Rx/ENROLLMENT FORM

THE INTAKE/REIMBURSEMENT SPECIALIST ENTERS THE PATIENT AND PHYSICIAN INFO INTO CHiPS

IS THE INFO COMPLETE?

NO

THE INTAKE/REIMBURSEMENT SPECIALIST CONTACTS MD TO VERIFY RECEIPT & ACCURACY OF THE PATIENT'S Rx & THIS CONTACT IS RECORDED IN CHiPS

YES

AN INTAKE REPRESENTATIVE WILL MAKE 1 ATTEMPT TO REACH THE MD TO OBTAIN THE MISSING INFORMATION

THE MISING INFORMATION HAS NOT BEEN OBTAINED WITHIN 24 HOURS, THE Rx/ENROLLMENT FORM IS FAXED BACK TO THE MD WITH A REJECTION EXPLANATION LETTER

A NOTE IS ENTERED IN CHiPS THAT THE APPLICATION WAS REJECTED

THE INTAKE/REIMBURSEMENT SPECIALIST SENDS CONSENT FORM AND A COVER LETTER TO THE PATIENT

THE INTAKE/REIMBURSEMENT SPECIALIST FAXES A STATEMENT OF MEDICAL NECESSITY TO THE MD TO COMPLETE

THE PATIENT IS INFORMED OF THE COST OF THE PRODUCT AND IS GIVEN PAYMENT OPTIONS

ONCE PAYMENT IS RECEIVED, INTAKE/REIMBURSEMENT SUBMITS A COVERAGE APPROVAL FORM (STAPLED TO THE "COPY" OF THE Rx/ENROLLMENT FORM) TO THE PHARMACY TEAM AS NOTIFICATION TO PROCESS THE PATIENT'S Rx

FIG. 2A
THE PATIENT IS SHIPPED A XYREM SUCCESS PACKET VIA 2-DAY USPS.

THE PHARMACIST HOLDS THE PATIENT'S Rx UNTIL RECEIVING A COVERAGE APPROVAL FORM FROM INTAKE/REIMBURSEMENT.

THE PHYSICIAN IS INDICATED AS APPROVED IN THE PHYSICIAN SCREEN IN CHiPS.

THE PHARMACIST HOLDS THE PATIENT'S Rx UNTIL RECEIVING A COVERAGE APPROVAL FORM FROM INTAKE/REIMBURSEMENT.

IF ANY DISCIPLINARY ACTIONS ARE IDENTIFIED, THE DIR. OF PHARM. IS NOTIFIED AND TAKES ONE OF THE FOLLOWING ACTIONS:

- Rx IS PROCESSED AND THE PHARMACIST MONITORS THE MD.'S PROGRAM ACTIVITY
- Rx IS NOT PROCESSED AND THE MD IS NOTED AS UNAPPROVED

THE MD IS CONTACTED BY A PHARMACIST & INFORMED THAT THE PATIENT'S Rx CANNOT BE PROCESSED.

A LETTER IS SENT TO THE PATIENT INDICATING THAT THE Rx CANNOT BE PROCESSED & THE PATIENT IS INSTRUCTED TO CONTACT HIS/HER PHYSICIAN.

FIG. 2B
UPON RECEIPT OF THE COVERAGE APPROVAL FORM FROM INTAKE/REIMBURSEMENT, THE PHARMACY TECHNICIAN CONTACTS THE PATIENT TO COMPLETE THE "TECHNICIAN" SPECIFIED SECTIONS OF THE PATIENT COUNSELING CHECKLIST.

WERE THE PROGRAM MATERIALS READ?


THE PHARMACIST SCHEDULE THE PATIENT'S SHIPMENT FOR THE NEXT BUSINESS DAY OR THE NEXT BUSINESS DAY THE PATIENT IS AVAILABLE TO SIGN FOR THE PACKAGE.

THE PHARMACIST ENTERS THE Rx ORDER IN CHiPS, CREATING AN ORDER NUMBER.

THE PHARMACIST VERIFIES THE Rx AND ATTACHES THE VERIFICATION LABEL TO THE HARD COPY Rx.

A PICK TICKET IS GENERATED FOR THE ORDER & THE ORDER IS FORWARDED TO THE PHARMACY FOR FULFILLMENT.

THE SHIPMENT IS CONFIRMED IN CHiPS.

THE ORIGINAL Rx IS FILED WITH THE PHARMACY Rx'S IN NUMERICAL ORDER.

THE ORDER IS SHIPPED BY USPS EXPRESS MAIL.

THE RECEIPT OF THE MATERIALS IS CONFIRMED AND ANOTHER CALL IS SCHEDULED TO COUNSEL THE PATIENT BEFORE THE XYREM IS SHIPPED.

THE SHIPMENT MUST BE SENT TO THE PATIENT'S HOME ADDRESS UNLESS THE PATIENT IS TRAVELING OR HAS MOVED, IN WHICH CASE THE PHARMACIST WILL DETERMINE IF AN EXCEPTION WILL BE MADE. THE PATIENT OR THE PATIENT'S DESIGNEE MUST SIGN FOR THE PACKAGE UPON DELIVERY.

THE ORIGINAL Rx IS FILED WITH THE PHARMACY Rx'S IN NUMERICAL ORDER.

ALL XYREM INVENTORY IS CYCLE COUNTED AND RECONCILED WITH THE CHiPS SYSTEM QUANTITIES BEFORE THE DAYS SHIPMENTS ARE SENT OUT.

FIG. 2C
PHYSICIAN SUCCESS PROGRAM MATERIALS REQUEST

MD CALLS XYREM SUCCESS PROGRAM AT 1-866-XYREM88 TO REQUEST PROGRAM MATERIALS

MD DEMOGRAPHICS, DEA# AND DATE OF REQUEST ARE ENTERED INTO CHIPS

SDS REQUESTS MATERIALS TO BE SHIPPED TO THE MD VIA THE OMI FULFILLMENT WEBSITE AT WWW.PRAFULFILLMENT.COM

END

FIG. 3
Each week a refill report is generated of patients with 10 days of product remaining.

A copy of the refill report is provided to intake/reimbursement.

No sooner than 8 days before medication depletion, a pharmacy technician contacts the patient to complete the pre-delivery checklist.

Is the patient reached?

Yes → The pharmacy technician schedules the patient's shipment for the next business day or the next business day the patient is available to sign for the package.

The pharmacist enters the Rx order in CHiPS, creating an order number.

The pharmacist verifies the Rx and attaches the verification label to the hard copy Rx.

The shipment is confirmed in CHiPS.

A pick ticket is generated for the order & the order is forwarded to the pharmacy for fulfillment.

No → A message is left only mentioning SDS & a return number.

A note is entered in the CHiPS indicating the date the message was left for the patient.

All XYREM inventory is cycle counted and reconciled with the CHiPS system quantities before the days shipments are sent out.

Fig. 4A
THE PATIENT CALLS TO REQUEST AN EARLY REFILL

A NOTE CODE IS ENTERED IN CHIPS ON THE PATIENT SCREEN INDICATING THE EARLY REFILL REQUEST

THE PHARMACIST EVALUATES THE PATIENT'S COMPLIANCE WITH THERAPY AND/OR POSSIBLE PRODUCT DIVERSION, MISUSE OR OVERSE

THE PHARMACIST CONTACTS THE PRESCRIBING PHYSICIAN TO ALERT OF THE SITUATION AND CONFIRM IF THE PHYSICIAN APPROVES OF THE EARLY REFILL

A XYREM PROBLEM IDENTIFICATION & MANAGEMENT RISK DIVERSION REPORT IS COMPLETED & DOCUMENTED IN CHIPS. THE REPORT IS THEN FAXED TO OMI & THE ORIGINAL IS FILED IN A MONTHLY BATCH FILE

THE PATIENT MUST WAIT UNTIL THE NEXT SCHEDULED REFILL DATE TO RECEIVE ADDITIONAL PRODUCT

END

THE PATIENT MUST WAIT UNTIL THE NEXT SCHEDULED REFILL DATE TO RECEIVE ADDITIONAL PRODUCT

IS THE PATIENT WILL TO PAY?

THE PATIENT IS INFORMED OF THE COST OF THE PRODUCT AND IS GIVEN PAYMENT OPTIONS

ONCE PAYMENT IS RECEIVED THE ORDER IS RELEASED

INTAKE/REIMBURSEMENT SUBMITS A COVERAGE APPROVAL FORM TO THE PHARMACY TEAM AS NOTIFICATION THAT THE PATIENT'S REFILL REQUEST CAN BE PROCESSED

INTAKE/REIMBURSEMENT SUBMITS A COVERAGE APPROVAL FORM TO THE PHARMACY TEAM AS NOTIFICATION THAT THE PATIENT'S REFILL REQUEST CAN BE PROCESSED

THE PHARMACY TECHNICIAN CONTACTS THE PATIENT TO SCHEDULE SHIPMENT OF THE PRODUCT FOR THE NEXT BUSINESS DAY OR THE NEXT BUSINESS DAY THE PATIENT IS AVAILABLE TO SIGN FOR THE PACKAGE

THE PHARMACY TECHNICIAN CONTACTS THE PATIENT TO SCHEDULE SHIPMENT OF THE PRODUCT FOR THE NEXT BUSINESS DAY OR THE NEXT BUSINESS DAY THE PATIENT IS AVAILABLE TO SIGN FOR THE PACKAGE

CONTINUE

CONTINUE

FIG. 4B
UPON DETERMINING THAT A PATIENT IS UNINSURED OR UNDERINSURED, A REIMBURSEMENT SPECIALIST EXPLAINS THE NORD PROGRAM TO THE PATIENT AND FAXES AN APPLICATION REQUEST FORM TO NORD FOR THE PATIENT.

THE INTAKE/REIMBURSEMENT SPECIALIST DOCUMENTS IN CHIPS THAT AN APPLICATION HAS BEEN REQUESTED THROUGH NORD.

NORD MAILS A PAP APPLICATION TO THE PATIENT WITHIN 1 BUSINESS DAY.

IS THE PATIENT APPROVED?

NORD SENDS AN ACCEPTANCE LETTER TO THE PATIENT AND FAXES A VOUCHER TO SDS INDICATING THE PATIENT'S APPROVAL FOR THE PROGRAM.

AN INTAKE/REIMBURSEMENT SPECIALIST SUBMITS A COVERAGE APPROVAL FORM TO THE PHARMACY TEAM AS NOTIFICATION THAT THE PATIENT HAS BEEN APPROVED FOR COVERAGE THROUGH NORD.

NORD SENDS A DENIAL LETTER TO THE PATIENT.

SDS DOCUMENTS IN CHIPS THAT THE PATIENT WAS DENIED BY NORD.

FIG. 5
EACH WEEK, THE DIRECTOR OF PHARMACY TRANSFERS INVENTORY FOR THE WEEK'S SHIPMENTS TO A SEGREGATED WAREHOUSE LOCATION FOR PRODUCTION INVENTORY

A PURCHASE ORDER IS GENERATED FOR THE INVENTORY TRANSFERRED TO THE PRODUCTION LOCATION & IS FAXED TO THE OMI CONTROLLER

THE OMI CONTROLLER INVOICES SDS FOR THE PRODUCT MOVED TO PRODUCTION

FIG. 6

FIG. 7

FIG. 8
**PRESCRIPTION AND ENROLLMENT FORM**

**PRESCRIBER INFORMATION**

<table>
<thead>
<tr>
<th>Precriber's Name:</th>
<th>Office Contact:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Street Address:</td>
<td></td>
</tr>
<tr>
<td>City:</td>
<td>State:</td>
</tr>
<tr>
<td>Phone:</td>
<td>Fax:</td>
</tr>
<tr>
<td>License Number:</td>
<td>DEA Number:</td>
</tr>
<tr>
<td>MD Specialty:</td>
<td></td>
</tr>
</tbody>
</table>

**PRESCRIPTION FORM**

<table>
<thead>
<tr>
<th>Patient Name:</th>
<th>SS#:</th>
<th>DOB:</th>
<th>Sex M / F</th>
</tr>
</thead>
<tbody>
<tr>
<td>Address:</td>
<td>City:</td>
<td>State:</td>
<td>ZIP:</td>
</tr>
</tbody>
</table>

Rx: XYREM ORAL SOLUTION (500 mg/mL) 180 ML BOTTLE QUANTITY: ___ MONTHS SUPPLY

SIG: Take ____caps p.o. diluted in 60 mL water at H.S. and then again 2 1/2 to 4 hours later

Refills (circle one): 0 1 2 (Maximum of 3 month supply)

Date: __/___/____

**PRESCRIBER'S SIGNATURE**

**PHYSICIAN DECLARATION** - PLEASE CHECK EACH BOX TO BE COMPLETED AT INITIAL PRESCRIPTION ONLY

- [ ] I HAVE READ THE MATERIALS IN THE XYREM PHYSICIAN SUCCESS PROGRAM
- [ ] I VERIFY THAT THE PATIENT HAS BEEN EDUCATED WITH RESPECT TO XYREM PREPARATION, DOSING AND SCHEDULING
- [ ] I UNDERSTAND THAT XYREM IS APPROVED FOR THE TREATMENT OF CATAPLEXY IN PATIENTS WITH NARCOLESPY, AND THAT SAFETY OR EFFICACY HAS NOT BEEN ESTABLISHED FOR ANY OTHER INDICATION
- [ ] I UNDERSTAND THAT THE SAFETY OF DOSES GREATER THAN 9gm/DAY HAS NOT BEEN ESTABLISHED

**PATIENT INFORMATION**

<table>
<thead>
<tr>
<th>Best time to contact patient:</th>
<th>Day:</th>
<th>Night:</th>
<th>Evening:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Insurance Company Name:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Insured's Name:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Identification Number:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prescription Card:</td>
<td>No:</td>
<td>Yes:</td>
<td></td>
</tr>
<tr>
<td>if Yes, Carrier:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Policy/Group Number:</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Please attach copies of patient's insurance cards

Fax completed form to XYREM success program (toll-free) 1-866-470-1744
For information, call the XYREM team (toll free) at 1-866-XYREM88 (1-866-997-3688)

**FIG. 9**
PATIENT ASSISTANCE APPLICATION REQUEST FORM

DATE:

TO: PATIENT ASSISTANCE ORGANIZATION

FROM: SDS

FAX #: 203-798-2291

PLEASE SEND A XYREM PATIENT ASSISTANCE PROGRAM APPLICATION TO:

PATIENT NAME ___________________________________________________________

ADDRESS ________________________________________________________________

________________________________________________________

TELEPHONE: ( ) __________________________________________________________

PATIENT DOSAGE: _______ (GRAMS) TWICE NIGHTLY FOR A TOTAL DOSAGE OF _______ (GRAMS)

_______ BOTTLES (THREE MONTHS SUPPLY)

BACKGROUND INFORMATION:

_____________________________________________________________________

_____________________________________________________________________

_____________________________________________________________________

_____________________________________________________________________

_____________________________________________________________________

FIG. 10
SENSITIVE DRUG PHYSICIAN’S CERTIFICATE
OF MEDICAL NEED

PATIENT INFORMATION

DATE: ________________

NAME: ____________________________________________

LAST FIRST M

DATE OF BIRTH: __________________________

DRUG BEING PRESCRIBED: XYREM

DIAGNOSIS/CONDITION FOR WHICH DRUG IS BEING PRESCRIBED: __________________________

ICD-9: __________________________

PHYSICIAN INFORMATION

PHYSICIAN’S NAME (PLEASE PRINT): __________________________

PHYSICIAN’S SIGNATURE: __________________________ DATE: ________________

PLEASE FAX BACK TO SENSITIVE DRUG SUCCESS PROGRAM: (1-800-TOLL FREE NUMBER)

FIG. 12
# ACTIVITY REPORTS

<table>
<thead>
<tr>
<th>Report Category</th>
<th>Report Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SALES</strong></td>
<td></td>
</tr>
<tr>
<td>Rx BY ZIP (NEW AND TOTAL)</td>
<td>X X X</td>
</tr>
<tr>
<td>Rx BY PHYSICIAN BY ZIP</td>
<td></td>
</tr>
<tr>
<td>Rx BY ZIP</td>
<td>X X X</td>
</tr>
<tr>
<td>$ BY ZIP</td>
<td></td>
</tr>
<tr>
<td><strong>REGULATORY</strong></td>
<td></td>
</tr>
<tr>
<td># OF PHYSICIAN REGISTRIES</td>
<td>X</td>
</tr>
<tr>
<td># OF DENIED PHYSICIAN REGISTRIES AND REASON</td>
<td>X</td>
</tr>
<tr>
<td># OF COMPLETED PATIENT REGISTRIES</td>
<td>X</td>
</tr>
<tr>
<td># OF PROBLEM IDENTIFICATION &amp; MANAGEMENT RISK DIVERSION REPORTS COMPLETED</td>
<td>X</td>
</tr>
<tr>
<td># OF CYCLE COUNTS PERFORMED &amp; ACCURACY OF EACH</td>
<td>X</td>
</tr>
<tr>
<td><strong>QUALITY ASSURANCE</strong></td>
<td></td>
</tr>
<tr>
<td># OF PRODUCT DEFECTS/COMPLAINTS REPORTED, TYPE AND LOT #</td>
<td>X</td>
</tr>
<tr>
<td><strong>CALL CENTER</strong></td>
<td></td>
</tr>
<tr>
<td># OF CALLS RECEIVED</td>
<td>X</td>
</tr>
<tr>
<td># OF CALLS INITIATED</td>
<td>X</td>
</tr>
<tr>
<td># OF CALLS ANSWERED IN 30 SECONDS, ETC.</td>
<td>X</td>
</tr>
<tr>
<td>PERCENTAGE OF CALLS ANSWERED IN 30 SECONDS</td>
<td>X</td>
</tr>
<tr>
<td># OF ABANDONED CALLS</td>
<td>X</td>
</tr>
<tr>
<td>% OF ABANDONED CALLS</td>
<td>X</td>
</tr>
<tr>
<td>AVERAGE CALL LENGTH</td>
<td>X</td>
</tr>
<tr>
<td><strong>PHARMACY</strong></td>
<td></td>
</tr>
<tr>
<td># OF FAXED Rx/ENROLLMENT FORMS</td>
<td>X</td>
</tr>
<tr>
<td># OF MAILED Rx/ENROLLMENT FORMS</td>
<td>X</td>
</tr>
<tr>
<td># OF RxS SHIPPED WIN 1, 2, 3, 4 ETC. DAYS (FROM THE TIME INITIAL RECEIPT TO SHIPMENT OF Rx)</td>
<td>X</td>
</tr>
<tr>
<td># OF PATIENT SUCCESS PACKETS SHIPPED</td>
<td>X</td>
</tr>
</tbody>
</table>

**FIG. 13A**
<table>
<thead>
<tr>
<th>ACTIVITY REPORTS</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>PHARMACY</td>
<td>X</td>
</tr>
<tr>
<td># OF PHYSICIAN SUCCESS PACKETS SHIPPED</td>
<td>X</td>
</tr>
<tr>
<td># OF COMPLETED SHIPMENTS</td>
<td>X</td>
</tr>
<tr>
<td># OF INCOMPLETE SHIPMENTS AND REASON</td>
<td>X</td>
</tr>
<tr>
<td># OF SHIPPING ERRORS</td>
<td>X</td>
</tr>
<tr>
<td># OF PAP SHIPMENTS</td>
<td>X</td>
</tr>
<tr>
<td># OF PAP APPLICATIONS</td>
<td>X</td>
</tr>
<tr>
<td># OF PAP APPROVALS</td>
<td>X</td>
</tr>
<tr>
<td># OF CANCELED ORDERS</td>
<td>X</td>
</tr>
<tr>
<td># OF USPS ERRORS</td>
<td>X</td>
</tr>
<tr>
<td>INVENTORY</td>
<td>X</td>
</tr>
<tr>
<td># OF RETURNED PRODUCTS AND REASON</td>
<td>X</td>
</tr>
<tr>
<td># OF OUTDATED BOTTLES OF PRODUCT</td>
<td>X</td>
</tr>
<tr>
<td>INVENTORY COUNTS OF CONSIGNMENT &amp; PRODUCTION INVENTORY</td>
<td>X</td>
</tr>
<tr>
<td># OF UNITS RECEIVED</td>
<td>X</td>
</tr>
<tr>
<td>LOTS RECEIVED</td>
<td>X</td>
</tr>
<tr>
<td>REIMBURSEMENT</td>
<td>X</td>
</tr>
<tr>
<td># OF PENDED AND WHY</td>
<td>X</td>
</tr>
<tr>
<td># OF APPROVALS</td>
<td>X</td>
</tr>
<tr>
<td># OF DENIALS</td>
<td>X</td>
</tr>
<tr>
<td># OF REJECTIONS</td>
<td>X</td>
</tr>
<tr>
<td>PAYOR TYPES</td>
<td>X</td>
</tr>
</tbody>
</table>

**FIG. 13B**
<table>
<thead>
<tr>
<th>ACTIVITY REPORTS</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>PATIENT CARE</td>
<td>X</td>
</tr>
<tr>
<td># OF ADVERSE EVENTS REPORTED AND TYPE</td>
<td>X</td>
</tr>
<tr>
<td># OF ADVERSE EVENTS SENT TO OMI</td>
<td>X</td>
</tr>
<tr>
<td># OF DOSING PROBLEMS AND TYPE</td>
<td>X</td>
</tr>
<tr>
<td># OF NONCOMPLIANCE EPISODES AND REASON</td>
<td>X</td>
</tr>
<tr>
<td># OF PATIENT COUNSELED AND REASON</td>
<td>X</td>
</tr>
<tr>
<td># OF PATIENTS DISCONTINUED AND REASON</td>
<td>X</td>
</tr>
<tr>
<td>PATIENT CARE</td>
<td>X</td>
</tr>
<tr>
<td># OF PATIENTS REFERRED TO PHYSICIAN AND REASON</td>
<td>X</td>
</tr>
<tr>
<td># OF ACTIVE PATIENTS</td>
<td>X</td>
</tr>
<tr>
<td># OF NEW PATIENTS</td>
<td>X</td>
</tr>
<tr>
<td># OF RESTART PATIENTS</td>
<td>X</td>
</tr>
<tr>
<td># OF DISCONTINUED PATIENTS AND REASON</td>
<td>X</td>
</tr>
<tr>
<td>DRUG INFORMATION</td>
<td>X</td>
</tr>
<tr>
<td># OF DRUG INFORMATION REQUESTS AND TYPE</td>
<td>X</td>
</tr>
<tr>
<td># OF CALLS TRIAGED TO OMI</td>
<td>X</td>
</tr>
</tbody>
</table>

**FIG. 13C**
SENITIVE DRUG DISTRIBUTION SYSTEM AND METHOD

RELATED APPLICATION

This application is a divisional application of U.S. patent application Ser. No. 10/322,348, filed Dec. 17, 2002 now U.S. Pat. No. 7,668,730, which application is incorporated herein by reference.

FIELD OF THE INVENTION

The present invention relates to distribution of drugs, and in particular to the distribution of sensitive drugs.

BACKGROUND OF THE INVENTION

Sensitive drugs are controlled to minimize risk and ensure that they are not abused, or cause adverse reactions. Such sensitive drugs are approved for specific uses by the Food and Drug Administration, and must be prescribed by a licensed physician in order to be purchased by consumers. Some drugs, such as cocaine and other common street drugs are the object of abuse and illegal schemes to distribute for profit. Some schemes include Dr. shopping, diversion, and pharmacy thefts. A locked cabinet or safe is a requirement for distribution of some drugs.

Certain agents, such as gamma hydroxy buterate (GHB) are also abused, yet also are effective for therapeutic purposes such as treatment of daytime cataplexy in patients with narcolepsy. Some patients however, will obtain prescriptions from multiple doctors, and have them filled at different pharmacies. Still further, an unscrupulous physician may actually write multiple prescriptions for a patient, or multiple patients, who use cash to pay for the drugs. These patients will then sell the drug to dealers or others for profit.

There is a need for a distribution system and method that directly addresses these abuses. There is a further need for such a system and method that provides education and limits the potential for such abuse.

SUMMARY OF THE INVENTION

A drug distribution system and method utilizes a central pharmacy and database to track all prescriptions for a sensitive drug. Information is kept in a central database regarding all physicians allowed to prescribe the sensitive drug, and all patients receiving the drug. Abuses are identified by monitoring data in the database for prescription patterns by physicians and prescriptions obtained by patients. Further verification is made that the physician is eligible to prescribe the drug by consulting a separate database for a valid DEA license, and optionally state medical boards to determine whether any corrective or approved disciplinary actions relating to controlled substances have been brought against the physician. Multiple controls beyond those for traditional drugs are imposed on the distribution depending on the sensitivity of the drug.

Education is provided to both physician and patient. Prior to shipping the drug for the first time, the patient is contacted to ensure that product and abuse related educational materials have been received and/or read. The patient may provide the name of a designee to the central pharmacy who is authorized to accept shipment of the drug. Receipt of the initial drug shipment is confirmed by contacting the patient. Either a phone call or other communication to the patient within a set time after delivery may be made to ensure receipt. Further, a courier service’s tracking system is used to confirm delivery in further embodiments. If a shipment is lost, an investigation is launched to find it.

In one embodiment, the drug may be shipped by the central pharmacy to another pharmacy for patient pick-up. The second pharmacy’s ability to protect against diversion before shipping the drug must be confirmed. This ability may be checked through NTIS and State Boards of Pharmacy.

Prescription refills are permitted in the number specified in the original prescription. In addition, if a prescription refill is requested by the patient prior to the anticipated due date, such refills will be questioned. A lost, stolen, destroyed or spilled prescription/supply is documented and replaced to the extent necessary to honor the prescription, and will also cause a review or full investigation.

The exclusive central database contains all relevant data related to distribution of the drug and process of distributing it, including patient, physician and prescription information. Several queries and reports are run against the database to provide information which might reveal potential abuse of the sensitive drug, such as early refills.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a block diagram of a computer system for use in implementing the system and method of the present invention.

FIGS. 2A, 2B and 2C are a flowchart describing a method for sensitive drug distribution at least partially utilizing a computer system such as that shown in FIG. 1.

FIG. 3 is a flowchart of a physician success program at least partially implemented on a computer system such as that shown in FIG. 1.

FIGS. 4A and 4B are a flowchart describing a method for handling refill requests at least partially utilizing a computer system such as that shown in FIG. 1.

FIG. 5 is a flowchart of a process for requesting special reimbursement when a patient is uninsured or underinsured at least partially utilizing a computer system as that shown in FIG. 1.

FIG. 6 is a flowchart of a process for inventory control at least partially utilizing a computer system such as that shown in FIG. 1.

FIG. 7 is a block diagram of database fields.

FIG. 8 is a block diagram showing a list of queries against the database fields.

FIG. 9 is a copy of one example prescription and enrollment form.

FIG. 10 is a copy of one example of a NORD application request form for patient financial assistance.

FIG. 11 is a copy of one example voucher request for medication for use with the NORD application request form of FIG. 10.

FIG. 12 is a copy of certificate of medical need.

FIGS. 13A, 13B and 13C are descriptions of sample reports obtained by querying a central database having fields represented in FIG. 7.

DETAILED DESCRIPTION OF THE INVENTION

In the following description, reference is made to the accompanying drawings that form a part hereof, and in which is shown by way of illustration specific embodiments in which the invention may be practiced. These embodiments are described in sufficient detail to enable those skilled in the art to practice the invention, and it is to be understood that other embodiments may be utilized and that structural, logical
and electrical changes may be made without departing from the scope of the present invention. The following description is, therefore, not to be taken in a limited sense, and the scope of the present invention is defined by the appended claims.

The functions or algorithms described herein are implemented in software or a combination of software and human implemented procedures in one embodiment. The software comprises computer executable instructions stored on computer readable media such as memory or other type of storage devices. The term “computer readable media” is also used to represent carrier waves on which the software is transmitted. Further, such functions correspond to modules, which are software, hardware, firmware of any combination thereof. Multiple functions are performed in one or more modules as desired, and the embodiments described are merely examples. The software is executed on a digital signal processor, ASIC, microprocessor, or other type of processor operating on a computer system, such as a personal computer, server or other computer system.

A sensitive drug is one which can be abused, or has addiction properties or other properties that render the drug sensitive. One example of such a drug is sodium oxybate, also known as gamma hydroxy butyrate (GHB C₂H₇NaO₃) which is useful for treatment of cataplexy in patients with narcolepsy. GHB is marketed under the trademark of Xyrem® (sodium oxybate oral solution), which trademark can be used interchangeably with GHB herein. Sensitive drugs also include narcotics or other drugs which require controls on their distribution and use to monitor behaviors to prevent abuse and adverse side effects.

In one embodiment, Xyrem® is subject to a restricted distribution program. One aspect of the program is to educate physicians and patients about the risks and benefits of Xyrem, including support via ongoing contact with patients and a toll free helpline. Initial prescriptions are filled only after a prescriber and patient have received and read the educational materials. Further, patient and prescribing physician registries are maintained and monitored to ensure proper distribution.

In a further embodiment, bulk sodium oxybate is manufactured at a single site, as is the finished drug product. Following manufacture of the drug product, it is stored at a facility compliant with FDA Schedule III regulations, where a consignment inventory is maintained. The inventory is owned by a company, and is managed by a central pharmacy, which maintains the consignment inventory. Xyrem® is distributed and dispensed through a primary and exclusive central pharmacy, and is not stocked in retail pharmacy outlets.

FIG. 1 is a simplified block diagram of a computer system 100, such as a personal computer for implementing at least a portion of the methods described herein. A central processing unit (CPU) 110 executes computer programs stored on a memory 120. Memory 120 in one embodiment comprises one or more levels of cache as desired to speed execution of the program and access to data on which the programs operate. The CPU is directly coupled to memory 120 in one embodiment. Both CPU 110 and memory 120 are coupled to a bus 130. A storage 140, I/O 150 and communications 160 are also coupled to the bus 130. Storage 140 is usually a long term storage device, such as a disk drive, tape drive, DVD, CD or other type of storage device. In one embodiment, storage 140 is used to house a database for use with the present invention. I/O 150 comprises keyboards, sound devices, displays and other mechanisms by which a user interacts with the computer system 100. Communications 160 comprises a network, phone connection, local area network, wide area network or other mechanism for communicating with external devices. Such external devices comprise servers, other peer computers and other devices. In one embodiment, such external device comprises a database server that is used in place of the database on storage 140. Other computer system architectures capable of executing software and interacting with a database and users may also be used. Appropriate security measures such as encryption are used to ensure confidentiality. Further, data integrity and backup measures are also used to prevent data loss.

FIGS. 2A, 2B and 2C represent an initial prescription order entry process for a sensitive drug, such as Xyrem. At 202, a medical doctor (MD) sends a RX/enrollment form via mail, fax, email or other means to an intake/reimbursement specialist at 204, who makes a copy of the RX/enrollment form that is stamped “copy”. The original fax is forwarded to a pharmacy team. The enrollment form contains prescriber information, prescription information, checkboxes for the prescriber indicating they have read materials, educated the patient, understand the use in treatment, and understand certain safety information, and also contains patient information.

The prescriber information contains standard contact information as well as license number, DEA number and physician specialty. Patient and prescription information includes name, social security number, date of birth, gender, contact information, drug identification, patient’s appropriate dosage, and number of refills allowed, along with a line for the prescriber’s signature. Patient insurance information is also provided.

There are two workflows involved at the pharmacy team, intake reimbursement 206 and pharmacy workflow 208, which may proceed in parallel or serially. The intake workflow 206 starts with an intake reimbursement specialist entering the patient and physician information into an application/database referred to as CHIPS, which is used to maintain a record of a client home infusion program (CHIP) for Xyrem®. A check is made to ensure the information is complete at 212. If not, at 214, an intake representative attempts to reach the MD or prescriber to obtain the missing information. If the missing information has not been obtained within a predetermined period of time, such as 24 hours at 216, the RX/Enrollment form is sent back to the MD with a rejection explanation. A note is entered in CHIPS that the application was rejected.

If the information is complete at 212, the MD is contacted at 220 to verify receipt and accuracy of the patient’s Rx. This contact is recorded in CHIPS. The intake and reimbursement specialist then sends a consent form and a cover letter to the patient at 224. The insurance provider is contacted at 226 to verify coverage and benefits. At 228, a determination is made regarding coverage for the drug. If it is not available, it is determined at 230 whether the patient is willing and able to pay. If not, a process is performed for handling patients who are uninsured or underinsured. In one embodiment, the process is referred to as a NORD process.

If the patient is willing and able to pay at 230, the patient is informed of the cost of the product and is given payment options at 234. At 236, once payment is received, the intake reimbursement specialist submits a coverage approval form with the enrollment form to the pharmacy team as notification to process the patient’s prescription. If coverage is approved at 228, the intake reimbursement specialist also submits the coverage approval form with the enrollment form to the pharmacy team as notification to process the patient’s prescription. Processing of the prescription is described below.
Upon receipt and initial processing of the prescription enrollment form and sending an original to the pharmacy workflow block 208, the patient is shipped a Xyrem® success packet via mail. In one embodiment, the Xyrem® success packet contains educational material for a patient that advises of the proper use, care, and handling of the drug and consequences of diversion at 268. The medical doctor’s credentials are checked to determine if the physician has a current DEA license to prescribe controlled substances and if he or she has had any actions related to misuse/misprescribing of controlled drugs against him or her, within a predetermined time, such as three months at 270. If they have, a pharmacist holds the prescription until receiving a coverage approval form from the intake reimbursement specialist at 272.

If the credentials have not been recently checked, the pharmacist verifies the credentials and enters all findings in the database at 274. If the credentials are approved at 276, the physician is indicated as approved in a physician screen populated by information from the database at 280. The prescription is then held pending coverage approval at 282.

If any disciplinary actions are identified, as referenced at block 278, management of the pharmacy is notified and either approves processing of the prescription with continued monitoring of the physician, or processing of the prescription is not performed, and the physician is noted in the database as unapproved at 284. The enrollment form is then mailed back to the physician with a cover letter reiterating that the prescription cannot be processed at 288. The patient is also sent a letter at 290 indicating that the prescription cannot be processed and the patient is instructed to contact their physician.

Actual filling of the approved prescription begins with receipt of the coverage approval form as indicated at 240. The patient is contacted by the pharmacy, such as by a technician to complete a technician section of a patient counseling checklist. If a pharmacist verifies that the program materials were not read at 242, the receipt of the material is confirmed at 244 and another call is scheduled to counsel the patient before the drug is shipped.

If the program materials were read at 242, the checklist is completed at 246 and the technician transfers the patient to the pharmacist who reviews the entire checklist and completes remaining pharmacist specified sections. At 248, the pharmacist indicates in the database that the patient counseling and checklist was successfully completed, indicating the date completed.

At 250, the pharmacist schedules the patient’s shipment for the next business day or the next business day that the patient or designee is able to pick up the package. Further, as indicated at 252, the shipment must be sent to the patient’s home address unless the patient is traveling or has moved. In that event, the pharmacist may determine that an exception may be made. The patient or the patient’s designee who is at least 18 years old, must sign for the package upon delivery.

At 254, the pharmacist enters the prescription order in the database, creating an order number. The pharmacist then verifies at 256 the prescription and attaches a verification label to the hard copy prescription. At 258, a pick ticket is generated for the order and the order is forwarded to the pharmacy for fulfillment. The shipment is confirmed in the database at 260, and the order is shipped by USPS Express Mail. Use of the US mail invokes certain criminal penalties for unauthorized diversion. Optionaly, other mail services may be used. Potential changes in the law may also bring criminal penalties into play. Following shipment, the patient is called by the central pharmacy to confirm that the prescription was received.

As noted at 266, for the sensitive drug, Xyrem, all inventory is cycle counted and reconciled with the database system quantities before shipments for the day are sent. This provides a very precise control of the inventory.

A physician success program materials request process begins at 310 in FIG. 3. At 320, the MD calls to the central pharmacy to request program materials. A special phone number is provided. MD demographics, DEA number, and data or request are entered into the database at 330. At 340, a request is made to ship the materials to the MD via a fulfillment website, or other mechanism. The request process ends at 350.

A refill request process begins at 302 in FIGS. 4A and 4B. There are two different paths for refills. A first path beginning at 404 involves generating a report from the central database of patients with a predetermined number of days or product remaining. A second path beginning at 406 is followed when a patient calls to request an early refill.

In the first path, a copy of the report is provided to an intake reimbursement specialist at 408. No sooner than 6 days before the medication depletion, a pharmacy technician contacts the patient at 410 to complete the pre-delivery checklist. At 412, if the patient is not reached, a message is left mentioning the depletion, and a return number at 414. A note is also entered into the database indicating the date the message was left at 416.

If the patient is reached at 412, the next shipment is scheduled at 418. The prescription is entered into the database creating an order at 420, the pharmacist verifies the prescription and attaches a verification label at 422 and the shipment is confirmed in the database at 424. Note at 426 that the inventory is cycle counted and reconciled with the database quantities before the shipments for a day or other time period are sent. A pick ticket is generated for the order and the order is forwarded for fulfillment at 428, with the first path ending at 430.

The second path, beginning at 406 results in a note code being entered into the database on a patient screen indicating an early refill request at 432. The pharmacist evaluates the patient’s compliance with therapy or possible product diversion, misuse or over-use at 436. In one embodiment, cash payers are also identified. The pharmacist then contacts the prescribing physician to alert them of the situation and confirm if the physician approves of the early refill at 438. If the physician does not approve as indicated at 440, the patient must wait until the next scheduled refill date to receive additional product as indicated at 442, and the process ends at 444.

If the physician approves at 440, the pharmacist enters a note in the database on a patient screen that the physician approves the request at 446. The pharmacist notifies an intake reimbursement specialist to contact the patient’s insurance provider to verify coverage for the early refill at 448. If the insurance provider will pay as determined at 450, the specialist submits the coverage approval form as notification that the refill may be processed at 452. At 454, the pharmacy technician contacts the patient to schedule shipment of the product for the next business day, and the process of filling the order is continued at 456 by following the process beginning at 240.

If the insurance provider will not pay at 450, it is determined whether the patient is willing and/or able to pay at 458. If not, the patient must wait until the next scheduled refill date to receive additional product at 460. If it was determined at 458 that the patient was willing and able to pay, the patient is informed of the cost of the product and is given payment options at 462. Once payment is received as indicated at 464, the specialist submits a coverage approval form to the pharmacy team as notification that the refill request can be pro-
process at 466. At 468, the pharmacy technician contacts the patient to schedule shipment. The process of filling the order is continued at 470 by following the process beginning at 240.

A process, referred to as a NORD process in one embodiment, is used to determine whether donated, third-party funds are available for paying for prescriptions where neither insurance will, nor the patient can pay. The process begins at 510 upon determining that a patient is uninsured or underinsured. A reimbursement specialist explains the NORD program to the patient and requests an application form. At 515, the intake reimbursement specialist documents the application in the database that an application has been received through NORD. At 520, NORD mails an application to the patient within one business day.

A determination is made at 525 by NORD whether the patient is approved. If not, at 530, NORD sends a denial letter to the patient, and it is documented in the database at 540 that the patient was denied by NORD. If the patient is approved, NORD sends an acceptance letter to the patient and faxes a voucher to the central pharmacy (SDS in one embodiment) to indicate the approval at 545. At 550, an intake reimbursement specialist submits a coverage approval form to the pharmacy team as notification that the patient has been approved for coverage. The process of filling the order is continued at 555 by following the process beginning at 240.

An inventory control process is illustrated in FIG. 6 beginning at 610. Each week, a responsible person at the central pharmacy, such as the director of the pharmacy transfers inventory for the week's shipments to a segregated warehouse location for production inventory. At 620, a purchase order is generated for the inventory transferred to the production location and is sent, such as by fax, to a controller, such as the controller of the company that obtained approval for distribution and use of the sensitive drug. At 630, the controller invoices the central pharmacy for the product moved to production. The process ends at 640.

The central database described above is a relational database running on the system of FIG. 1, or a server based system having a similar architecture coupled to workstations via a network, as represented by communications 160. The database is likely stored in storage 140, and contains multiple fields of information as indicated at 700 in FIG. 7. The organization and groupings of the fields are shown in one format for convenience. It is recognized that many different organizations or schemas may be utilized. In one embodiment, the groups of fields compriseprescriber fields 710, patient fields 720, prescription fields 730 and insurance fields 740. For purposes of illustration, all the entries described with respect to the above processes are included in the fields. In further embodiments, no such groupings are made, and the data is organized in a different manner.

Several queries are illustrated at 800 in FIG. 8. There may be many other queries as required by individual state reporting requirements. A first query at 810 is used to identify prescriptions written by physician. The queries may be written in structured query language, natural language queries or in any other manner compatible with the database. A second query 820 is used to pull information from the database related to prescriptions by patient name. A third query 830 is used to determine prescriptions by frequency, and a n-th query finds prescriptions by dose at 840. Using query languages combined with the depth of data in the central database allows many other methods of investigating for potential abuse of the drugs. The central database ensures that all prescriptions, prescribers and patients are tracked and subject to such investigations. In further embodiments, the central database may be distributed among multiple computers provided a query operates over all data relating to such prescriptions, prescribers and patients for the drug.

An example of one prescription and enrollment form is shown at 900 in FIG. 9. As previously indicated, several fields are included for prescriber information, prescription information and patient information. FIG. 10 is a copy of one example NORD application request form used to send the application to the patient for financial assistance. FIG. 11 is a copy of one example application for financial assistance as requested by form 1000. The form requires both patient and physician information. Social security number information is also required. The form provides information for approving the financial assistance and for tracking assistance provided.

FIG. 12 is a copy of one example voucher request for medication for use with the NORD application request form of FIG. 10. In addition to patient and physician information, prescription information and diagnosis information is also provided.

FIGS. 13A, 13B and 13C are descriptions of sample reports obtained by querying a central database having fields represented in FIG. 7. The activities grouped by sales, regulatory, quality assurance, call center, pharmacy, inventory, reimbursement, patient care and drug information. Each report has an associated frequency or frequencies. The reports are obtained by running queries against the database, with the queries written in one of many query languages.

While the invention has been described with respect to a Schedule III drug, it is useful for other sensitive drugs that are DEA or Federally scheduled drugs in Schedule II-V, as well as still other sensitive drugs where multiple controls are desired for distribution and use.

The invention claimed is:

1. A method of obtaining FDA (Food and Drug Administration) approval for a prescription drug, the method comprising:
   - maintaining with a computer processor an exclusive central computer database;
   - determining with the computer processor current and anticipated patterns of potential abuse of the prescription drug, wherein the anticipated patterns of potential abuse are determined in part by requiring entry of information into the exclusive central computer database such that any and all prescriptions for the prescription drug, for any and all patients being prescribed the prescription drug, and from any and all doctors allowed to prescribe the prescription drug, are processed only via the exclusive central computer database;
   - selecting with the computer processor multiple controls for distribution while maintaining the exclusive central computer database, the controls comprising controls beyond the controls for traditional prescription drugs based on the sensitivity of the prescription drug, the controls selected from the group consisting of communicating prescriptions from a physician to a pharmacy, identifying the physician's name, license and DEA (Drug Enforcement Agency) registration information, verifying the prescription; obtaining patient information, verifying the physician is eligible to prescribe the prescription drug by consulting the National Technical Information Services to determine whether the physician has an active DEA number and check on whether any actions are pending against the physician, providing comprehensive printed materials to the physician, contacting the patient's insurance company if any, verifying
patient registry information, providing comprehensive education information to the patient, verifying the patient has received or reviewed the educational materials, verifying the home address of the patient, shipping via US postal service or a commercial shipping service, receiving the name of an at least 18 year old designee to receive the sensitive drug after two attempts to deliver, launching an investigation when a shipment is lost, shipping to another pharmacy for delivery, requiring manufacture at a single location, releasing inventory in a controlled manner, questioning early refills, flagging repeat instances of lost, stolen, destroyed or spilled prescriptions, and making the database available to the DEA for checking for abuse patterns in data in the database, analyzing cash payments stored in the database, analyzing inappropriate questions stored in the database; displaying the selected controls to an output device; and receiving negotiation input from the FDA and adding with the computer processor further controls from the group until approval is obtained;

wherein the prescription drug is a scheduled drug in Schedules II-V of FDA subpart 4 regulation embodied in Title 21, CFR Part 314, existing on Apr. 1, 2005.

2. The method of claim 1 wherein initially selected controls comprise communicating prescriptions from a physician to a pharmacy, identifying the physician’s name, license and DEA registration information, verifying the prescription; obtaining patient information, verifying the physician is eligible to prescribe the prescription drug by consulting the National Technical Information Services to determine whether the physician has an active DEA number and check on whether any actions are pending against the physician, providing comprehensive education information to the patient, verifying the physician has an active DEA number and check on whether any actions are pending against the physician, providing comprehensive education information to the patient, verifying the home address of the patient, shipping via US postal service or a commercial shipping service, receiving the name of an at least 18 year old designee to receive the sensitive drug after two attempts to deliver, launching an investigation when a shipment is lost, shipping to another pharmacy for delivery, requiring manufacture at a single location, releasing inventory in a controlled manner, questioning early refills, flagging repeat instances of lost, stolen, destroyed or spilled prescriptions, and making the database available to the DEA for checking for abuse patterns in data in the database, analyzing cash payments stored in the database, analyzing inappropriate questions stored in the database;

displaying the selected controls to an output device; and receiving negotiation input from the FDA and adding with the computer processor further controls from the group until approval is obtained;

wherein the sensitive drug is a scheduled drug in Schedules II-IV of FDA subpart 4 regulation embodied in Title 21, CFR Part 314, wherein the Schedules II-IV are as of Apr. 1, 2005.

3. A method of obtaining FDA (Food and Drug Administration) approval for a sensitive drug, the method comprising: maintaining with a computer processor an exclusive central computer database;

determining with the computer processor current and anticipated patterns of potential abuse of the sensitive drug, wherein the anticipated patterns of potential abuse are determined in part by requiring entry of information into the exclusive central computer database such that any and all prescriptions for the sensitive drug, for any and all patients being prescribed the sensitive drug, and from any and all doctors allowed to prescribe the sensitive drug, are processed only via the exclusive central computer database;

selecting with the computer processor multiple controls for distribution while maintaining the exclusive central computer database, the controls comprising controls beyond the controls for traditional prescription drugs based on the sensitivity of the sensitive drug, the controls selected from the group consisting of communicating prescriptions from a physician to a pharmacy, identifying the physician’s name, license and DEA (Drug Enforcement Agency) registration information, verifying the prescription; obtaining patient information, verifying the physician is eligible to prescribe the prescription drug by consulting the National Technical Information Services to determine whether the physician has an active DEA number and check on whether any actions are pending against the physician, providing comprehensive printed materials to the physician, contacting the patient’s insurance company if any, verifying patient registry information, providing comprehensive education information to the patient, verifying the patient has received or reviewed the educational materials, verifying the home address of the patient, shipping via US postal service or a commercial shipping service, receiving the name of an at least 18 year old designee to receive the sensitive drug, confirming receipt of an initial shipment of the sensitive drug to the patient, returning the prescription drug after two attempts to deliver, launching an investigation when a shipment is lost, shipping to another pharmacy for delivery, requiring manufacture at a single location, releasing inventory in a controlled manner, questioning early refills, flagging repeat instances of lost, stolen, destroyed or spilled prescriptions, and making the database available to the DEA for checking for abuse patterns in data in the database, analyzing cash payments stored in the database, analyzing inappropriate questions stored in the database; displaying the selected controls to an output device; and receiving negotiation input from the FDA and adding with the computer processor further controls from the group until approval is obtained;

wherein the sensitive drug is a scheduled drug in Schedules II-IV of FDA subpart 4 regulation embodied in Title 21, CFR Part 314, wherein the Schedules II-IV are as of Apr. 1, 2005.

4. The method of claim 3, wherein initially selected controls comprise communicating prescriptions from a physician to a pharmacy, identifying the physician’s name, license and DEA registration information, verifying the prescription; obtaining patient information, verifying the physician is eligible to prescribe the sensitive drug by consulting the National Technical Information Services to determine whether the physician has an active DEA number and check on whether any actions are pending against the physician, providing comprehensive education information to the patient, verifying the patient has received or reviewed the educational materials, verifying the home address of the patient, shipping via US postal service or a commercial shipping service, receiving the name of an at least 18 year old designee to receive the sensitive drug, confirming receipt of an initial shipment of the sensitive drug to the patient, returning the prescription drug after two attempts to deliver, launching an investigation when a shipment is lost, shipping to another pharmacy for delivery, requiring manufacture at a single location, releasing inventory in a controlled manner, questioning early refills, flagging repeat instances of lost, stolen, destroyed or spilled prescriptions, and making the database available to the DEA for checking for abuse patterns in data.