	(Original Signature of Member)
	TH CONGRESS 1ST SESSION  H. R.
	amend the Federal Food, Drug, and Cosmetic Act with respect to human rug compounding and drug supply chain security, and for other purposes.
	IN THE HOUSE OF REPRESENTATIVES
Mr.	UPTON introduced the following bill; which was referred to the Committee on
	A BILL
То	amend the Federal Food, Drug, and Cosmetic Act with
	respect to human drug compounding and drug supply
	chain security, and for other purposes.
1	Be it enacted by the Senate and House of Representa-
2	tives of the United States of America in Congress assembled,
3	SECTION 1. SHORT TITLE.
4	This Act may be cited as the "Drug Quality and Se-
5	curity Act".
6	SEC. 2. REFERENCES IN ACT; TABLE OF CONTENTS.
7	(a) References in Act.—Except as otherwise spec-
8	ified, amendments made by this Act to a section or other

- 1 provision of law are amendments to such section or other
- 2 provision of the Federal Food, Drug, and Cosmetic Act
- 3 (21 U.S.C. 301 et seq.).
- 4 (b) Table of Contents of
- 5 this Act is as follows:
  - Sec. 1. Short title.
  - Sec. 2. References in Act; table of contents.

## TITLE I—DRUG COMPOUNDING

- Sec. 101. Short title.
- Sec. 102. Voluntary outsourcing facilities.
- Sec. 103. Penalties.
- Sec. 104. Regulations.
- Sec. 105. Enhanced communication.
- Sec. 106. Severability.
- Sec. 107. GAO study.

## TITLE II—DRUG SUPPLY CHAIN SECURITY

- Sec. 201. Short title.
- Sec. 202. Pharmaceutical distribution supply chain.
- Sec. 203. Enhanced drug distribution security.
- Sec. 204. National standards for prescription drug wholesale distributors.
- Sec. 205. National standards for third-party logistics providers; uniform national policy.
- Sec. 206. Penalties.
- Sec. 207. Conforming amendment.
- Sec. 208. Savings clause.

## 6 TITLE I—DRUG COMPOUNDING

- 7 SEC. 101. SHORT TITLE.
- 8 This Act may be cited as the "Compounding Quality
- 9 Act".
- 10 SEC. 102. VOLUNTARY OUTSOURCING FACILITIES.
- 11 (a) IN GENERAL.—Subchapter A of chapter V (21
- 12 U.S.C. 351 et seq.) is amended—
- 13 (1) by redesignating section 503B as section
- 14 503C; and

1	(2) by inserting after section 503A the fol-
2	lowing new section:
3	"SEC. 503B. OUTSOURCING FACILITIES.
4	"(a) In General.—Sections $502(f)(1)$ , $505$ , and $582$
5	shall not apply to a drug compounded by or under the
6	direct supervision of a licensed pharmacist in a facility
7	that elects to register as an outsourcing facility if each
8	of the following conditions is met:
9	"(1) REGISTRATION AND REPORTING.—The
10	drug is compounded in an outsourcing facility that
11	is in compliance with the requirements of subsection
12	(b).
13	"(2) Bulk drug substances.—The drug is
14	compounded in an outsourcing facility that does not
15	compound using bulk drug substances (as defined in
16	section 207.3(a)(4) of title 21, Code of Federal Reg-
17	ulations (or any successor regulation)), unless—
18	"(A)(i) the bulk drug substance appears on
19	a list established by the Secretary identifying
20	bulk drug substances for which there is a clin-
21	ical need, by—
22	"(I) publishing a notice in the Federal
23	Register proposing bulk drug substances to
24	be included on the list, including the ra-
25	tionale for such proposal;

1	"(II) providing a period of not less
2	than 60 calendar days for comment on the
3	notice; and
4	"(III) publishing a notice in the Fed-
5	eral Register designating bulk drug sub-
6	stances for inclusion on the list; or
7	"(ii) the drug compounded from such bulk
8	drug substance appears on the drug shortage
9	list in effect under section 506E at the time of
10	compounding, distribution, and dispensing;
11	"(B) if an applicable monograph exists
12	under the United States Pharmacopeia, the Na-
13	tional Formulary, or another compendium or
14	pharmacopeia recognized by the Secretary for
15	purposes of this paragraph, the bulk drug sub-
16	stances each comply with the monograph;
17	"(C) the bulk drug substances are each
18	manufactured by an establishment that is reg-
19	istered under section 510 (including a foreign
20	establishment that is registered under section
21	510(i)); and
22	"(D) the bulk drug substances are each ac-
23	companied by a valid certificate of analysis.
24	"(3) Ingredients (other than bulk drug
25	SUBSTANCES).—If any ingredients (other than bulk

1	drug substances) are used in compounding the drug,
2	such ingredients comply with the standards of the
3	applicable United States Pharmacopeia or National
4	Formulary monograph, if such monograph exists, or
5	of another compendium or pharmacopeia recognized
6	by the Secretary for purposes of this paragraph if
7	any.
8	"(4) Drugs withdrawn or removed be-
9	CAUSE UNSAFE OR NOT EFFECTIVE.—The drug does
10	not appear on a list published by the Secretary of
11	drugs that have been withdrawn or removed from
12	the market because such drugs or components of
13	such drugs have been found to be unsafe or not ef-
14	fective.
15	"(5) Essentially a copy of an approved
16	DRUG.—The drug is not essentially a copy of one or
17	more approved drugs.
18	"(6) Drugs presenting demonstrable dif-
19	FICULTIES FOR COMPOUNDING.—The drug—
20	"(A) is not identified (directly or as part
21	of a category of drugs) on a list published by
22	the Secretary, through the process described in
23	subsection (c), of drugs or categories of drugs
24	that present demonstrable difficulties for
25	compounding that are reasonably likely to lead

1	to an adverse effect on the safety or effective-
2	ness of the drug or category of drugs, taking
3	into account the risks and benefits to patients;
4	or
5	"(B) is compounded in accordance with all
6	applicable conditions identified on the list de-
7	scribed in subparagraph (A) as conditions that
8	are necessary to prevent the drug or category of
9	drugs from presenting the demonstrable dif-
10	ficulties described in subparagraph (A).
11	"(7) Elements to assure safe use.—In the
12	case of a drug that is compounded from a drug that
13	is the subject of a risk evaluation and mitigation
14	strategy approved with elements to assure safe use
15	pursuant to section 505-1, or from a bulk drug sub-
16	stance that is a component of such drug, the
17	outsourcing facility demonstrates to the Secretary
18	prior to beginning compounding that such facility
19	will utilize controls comparable to the controls appli-
20	cable under the relevant risk evaluation and mitiga-
21	tion strategy.
22	"(8) Prohibition on wholesaling.—The
23	drug will not be sold or transferred by an entity
24	other than the outsourcing facility that compounded
25	such drug. This paragraph does not prohibit admin-

1	istration of a drug in a health care setting or dis-
2	pensing a drug pursuant to a prescription executed
3	in accordance with section 503(b)(1).
4	"(9) Fees.—The drug is compounded in an
5	outsourcing facility that has paid all fees owed by
6	such facility pursuant to section 744K.
7	"(10) Labeling of drugs.—
8	"(A) Label.—The label of the drug in-
9	cludes—
10	"(i) the statement 'This is a com-
11	pounded drug.' or a reasonable comparable
12	alternative statement (as specified by the
13	Secretary) that prominently identifies the
14	drug as a compounded drug;
15	"(ii) the name, address, and phone
16	number of the applicable outsourcing facil-
17	ity; and
18	"(iii) with respect to the drug—
19	"(I) the lot or batch number;
20	"(II) the established name of the
21	drug;
22	"(III) the dosage form and
23	strength;
24	"(IV) the statement of quantity
25	or volume, as appropriate;

1	"(V) the date that the drug was
2	compounded;
3	"(VI) the expiration date;
4	"(VII) storage and handling in-
5	structions;
6	"(VIII) the National Drug Code
7	number, if available;
8	"(IX) the statement 'Not for re-
9	sale', and, if the drug is dispensed or
10	distributed other than pursuant to a
11	prescription for an individual identi-
12	fied patient, the statement 'Office Use
13	Only'; and
14	"(X) subject to subparagraph
15	(B)(i), a list of active and inactive in-
16	gredients, identified by established
17	name and the quantity or proportion
18	of each ingredient.
19	"(B) Container from
20	which the individual units of the drug are re-
21	moved for dispensing or for administration
22	(such as a plastic bag containing individual
23	product syringes) shall include—

1	"(i) the information described under
2	subparagraph (A)(iii)(X), if there is not
3	space on the label for such information;
4	"(ii) the following information to fa-
5	cilitate adverse event reporting:
6	www.fda.gov/medwatch and $1-800$ -FDA-
7	1088 (or any successor Internet Web site
8	or phone number); and
9	"(iii) directions for use, including, as
10	appropriate, dosage and administration.
11	"(C) Additional information.—The
12	label and labeling of the drug shall include any
13	other information as determined necessary and
14	specified in regulations promulgated by the Sec-
15	retary.
16	"(11) Outsourcing facility require-
17	MENT.—The drug is compounded in an outsourcing
18	facility in which the compounding of drugs occurs
19	only in accordance with this section.
20	"(b) REGISTRATION OF OUTSOURCING FACILITIES
21	AND REPORTING OF DRUGS.—
22	"(1) Registration of outsourcing facili-
23	TIES.—
24	"(A) ANNUAL REGISTRATION.—Upon
25	electing and in order to become an outsourcing

1	facility, and during the period beginning on Oc-
2	tober 1 and ending on December 31 of each
3	year thereafter, a facility—
4	"(i) shall register with the Secretary
5	its name, place of business, and unique fa-
6	cility identifier (which shall conform to the
7	requirements for the unique facility identi-
8	fier established under section 510), and a
9	point of contact email address; and
10	"(ii) shall indicate whether the
11	outsourcing facility intends to compound a
12	drug that appears on the list in effect
13	under section 506E during the subsequent
14	calendar year.
15	"(B) Availability of registration for
16	INSPECTION; LIST.—
17	"(i) Registrations.—The Secretary
18	shall make available for inspection, to any
19	person so requesting, any registration filed
20	pursuant to this paragraph.
21	"(ii) List.—The Secretary shall make
22	available on the public Internet Web site of
23	the Food and Drug Administration a list
24	of the name of each facility registered
25	under this subsection as an outsourcing fa-

1	cility, the State in which each such facility
2	is located, whether the facility compounds
3	from bulk drug substances, and whether
4	any such compounding from bulk drug
5	substances is for sterile or nonsterile
6	drugs.
7	"(2) Drug reporting by outsourcing fa-
8	CILITIES.—
9	"(A) In General.—Upon initially reg-
10	istering as an outsourcing facility, once during
11	the month of June of each year, and once dur-
12	ing the month of December of each year, each
13	outsourcing facility that registers with the Sec-
14	retary under paragraph (1) shall submit to the
15	Secretary a report—
16	"(i) identifying the drugs compounded
17	by such outsourcing facility during the pre-
18	vious 6-month period; and
19	"(ii) with respect to each drug identi-
20	fied under clause (i), providing the active
21	ingredient, the source of such active ingre-
22	dient, the National Drug Code number of
23	the source drug or bulk active ingredient,
24	if available, the strength of the active in-
25	gredient per unit, the dosage form and

1	route of administration, the package de-
2	scription, the number of individual units
3	produced, and the National Drug Code
4	number of the final product, if assigned.
5	"(B) FORM.—Each report under subpara-
6	graph (A) shall be prepared in such form and
7	manner as the Secretary may prescribe by regu-
8	lation or guidance.
9	"(C) Confidentiality.—Reports sub-
10	mitted under this paragraph shall be exempt
11	from inspection under paragraph (1)(B)(i), un-
12	less the Secretary finds that such an exemption
13	would be inconsistent with the protection of the
14	public health.
15	"(3) Electronic registration and report-
16	ING.—Registrations and drug reporting under this
17	subsection (including the submission of updated in-
18	formation) shall be submitted to the Secretary by
19	electronic means unless the Secretary grants a re-
20	quest for waiver of such requirement because use of
21	electronic means is not reasonable for the person re-
22	questing waiver.
23	"(4) Risk-based inspection frequency.—
24	"(A) In General.—Outsourcing facili-
25	ties—

1	"(i) shall be subject to inspection pur-
2	suant to section 704; and
3	"(ii) shall not be eligible for the ex-
4	emption under section 704(a)(2)(A).
5	"(B) RISK-BASED SCHEDULE.—The Sec-
6	retary, acting through one or more officers or
7	employees duly designated by the Secretary,
8	shall inspect outsourcing facilities in accordance
9	with a risk-based schedule established by the
10	Secretary.
11	"(C) RISK FACTORS.—In establishing the
12	risk-based schedule, the Secretary shall inspect
13	outsourcing facilities according to the known
14	safety risks of such outsourcing facilities, which
15	shall be based on the following factors:
16	"(i) The compliance history of the
17	outsourcing facility.
18	"(ii) The record, history, and nature
19	of recalls linked to the outsourcing facility.
20	"(iii) The inherent risk of the drugs
21	compounded at the outsourcing facility.
22	"(iv) The inspection frequency and
23	history of the outsourcing facility, includ-
24	ing whether the outsourcing facility has

1	been inspected pursuant to section 704
2	within the last 4 years.
3	"(v) Whether the outsourcing facility
4	has registered under this paragraph as an
5	entity that intends to compound a drug
6	that appears on the list in effect under sec-
7	tion 506E.
8	"(vi) Any other criteria deemed nec-
9	essary and appropriate by the Secretary
10	for purposes of allocating inspection re-
11	sources.
12	"(5) Adverse event reporting.—
13	Outsourcing facilities shall submit adverse event re-
14	ports to the Secretary in accordance with the con-
15	tent and format requirements established through
16	guidance or regulation under section 310.305 of title
17	21, Code of Federal Regulations (or any successor
18	regulations).
19	"(e) Regulations.—
20	"(1) In General.—The Secretary shall imple-
21	ment the list described in subsection (a)(6) through
22	regulations.
23	"(2) Advisory committee on
24	COMPOUNDING.—Before issuing regulations to im-
25	plement subsection (a)(6), the Secretary shall con-

1	vene and consult an advisory committee on
2	compounding. The advisory committee shall include
3	representatives from the National Association of
4	Boards of Pharmacy, the United States Pharma-
5	copeia, pharmacists with current experience and ex-
6	pertise in compounding, physicians with background
7	and knowledge in compounding, and patient and
8	public health advocacy organizations.
9	"(3) Interim list.—
10	"(A) In General.—Before the effective
11	date of the regulations finalized to implement
12	subsection (a)(6), the Secretary may designate
13	drugs, categories of drugs, or conditions as de-
14	scribed such subsection by—
15	"(i) publishing a notice of such sub-
16	stances, drugs, categories of drugs, or con-
17	ditions proposed for designation, including
18	the rationale for such designation, in the
19	Federal Register;
20	"(ii) providing a period of not less
21	than 60 calendar days for comment on the
22	notice; and
23	"(iii) publishing a notice in the Fed-
24	eral Register designating such drugs, cat-
25	egories of drugs, or conditions.

1	"(B) Sunset of Notice.—Any notice
2	provided under subparagraph (A) shall not be
3	effective after the earlier of—
4	"(i) the date that is 5 years after the
5	date of enactment of the Compounding
6	Quality Act; or
7	"(ii) the effective date of the final reg-
8	ulations issued to implement subsection
9	(a)(6).
10	"(4) UPDATES.—The Secretary shall review,
11	and update as necessary, the regulations containing
12	the lists of drugs, categories of drugs, or conditions
13	described in subsection (a)(6) regularly, but not less
14	than once every 4 years. Nothing in the previous
15	sentence prohibits submissions to the Secretary, be-
16	fore or during any 4-year period described in such
17	sentence, requesting updates to such lists.
18	"(d) Definitions.—In this section:
19	"(1) The term 'compounding' includes the com-
20	bining, admixing, mixing, diluting, pooling, reconsti-
21	tuting, or otherwise altering of a drug or bulk drug
22	substance to create a drug.
23	"(2) The term 'essentially a copy of an ap-
24	proved drug' means—

1	"(A) a drug that is identical or nearly
2	identical to an approved drug, or a marketed
3	drug not subject to section 503(b) and not sub-
4	ject to approval in an application submitted
5	under section 505, unless, in the case of an ap-
6	proved drug, the drug appears on the drug
7	shortage list in effect under section 506E at the
8	time of compounding, distribution, and dis-
9	pensing; or
10	"(B) a drug, a component of which is a
11	bulk drug substance that is a component of an
12	approved drug or a marketed drug that is not
13	subject to section 503(b) and not subject to ap-
14	proval in an application submitted under sec-
15	tion 505, unless there is a change that produces
16	for an individual patient a clinical difference, as
17	determined by the prescribing practitioner, be-
18	tween the compounded drug and the com-
19	parable approved drug.
20	"(3) The term 'approved drug' means a drug
21	that is approved under section 505 and does not ap-
22	pear on the list described in subsection (a)(4) of
23	drugs that have been withdrawn or removed from
24	the market because such drugs or components of

1	such drugs have been found to be unsafe or not ef-
2	fective.
3	"(4)(A) The term 'outsourcing facility' means a
4	facility at one geographic location or address that—
5	"(i) is engaged in the compounding of ster-
6	ile drugs;
7	"(ii) has elected to register as an
8	outsourcing facility; and
9	"(iii) complies with all of the requirements
10	of this section.
11	"(B) An outsourcing facility is not required to
12	be a licensed pharmacy.
13	"(C) An outsourcing facility may or may not
14	obtain prescriptions for identified individual pa-
15	tients.
16	"(5) The term 'sterile drug' means a drug that
17	is intended for parenteral administration, an oph-
18	thalmic or oral inhalation drug in aqueous format,
19	or a drug that is required to be sterile under Federal
20	or State law.".
21	"(d) Obligation to Pay Fees.—Payment of the fee
22	under section 744K, as described in subsection (a)(9),
23	shall not relieve an outsourcing facility that is licensed as
24	a pharmacy in any State that requires pharmacy licensing
25	fees of its obligation to pay such State fees.".

1	(b) Fees.—Subchapter C of chapter VII (21 U.S.C.
2	379f et seq.) is amended by adding at the end the fol-
3	lowing:
4	"PART 9—FEES RELATING TO OUTSOURCING
5	FACILITIES
6	"SEC. 744J. DEFINITIONS.
7	"In this part:
8	"(1) The term 'affiliate' has the meaning given
9	such term in section 735(11).
10	"(2) The term 'gross annual sales' means the
11	total worldwide gross annual sales, in United States
12	dollars, for an outsourcing facility, including the
13	sales of all the affiliates of the outsourcing facility.
14	"(3) The term 'outsourcing facility' has the
15	meaning given to such term in section 503B(d)(4).
16	"(4) The term 'reinspection' means, with re-
17	spect to an outsourcing facility, 1 or more inspec-
18	tions conducted under section 704 subsequent to an
19	inspection conducted under such provision which
20	identified noncompliance materially related to an ap-
21	plicable requirement of this Act, specifically to deter-
22	mine whether compliance has been achieved to the
23	Secretary's satisfaction.

1	"SEC. 744K. AUTHORITY TO ASSESS AND USE
2	OUTSOURCING FACILITY FEES.
3	"(a) Establishment and Reinspection Fees.—
4	"(1) In general.—For fiscal year 2015 and
5	each subsequent fiscal year, the Secretary shall, in
6	accordance with this subsection, assess and collect—
7	"(A) an annual establishment fee from
8	each outsourcing facility; and
9	"(B) a reinspection fee from each
10	outsourcing facility subject to a reinspection in
11	such fiscal year.
12	"(2) Multiple reinspections.—An
13	outsourcing facility subject to multiple reinspections
14	in a fiscal year shall be subject to a reinspection fee
15	for each reinspection.
16	"(b) Establishment and Reinspection Fee Set-
17	TING.—The Secretary shall—
18	"(1) establish the amount of the establishment
19	fee and reinspection fee to be collected under this
20	section for each fiscal year based on the method-
21	ology described in subsection (c); and
22	"(2) publish such fee amounts in a Federal
23	Register notice not later than 60 calendar days be-
24	fore the start of each such year.
25	"(c) Amount of Establishment Fee and Rein-
26	SPECTION FEE.—

1	"(1) In general.—For each outsourcing facil-
2	ity in a fiscal year—
3	"(A) except as provided in paragraph (4),
4	the amount of the annual establishment fee
5	under subsection (b) shall be equal to the sum
6	of—
7	"(i) \$15,000, multiplied by the infla-
8	tion adjustment factor described in para-
9	graph (2); plus
10	"(ii) the small business adjustment
11	factor described in paragraph (3); and
12	"(B) the amount of any reinspection fee (if
13	applicable) under subsection (b) shall be equal
14	to \$15,000, multiplied by the inflation adjust-
15	ment factor described in paragraph (2).
16	"(2) Inflation adjustment factor.—
17	"(A) In general.—For fiscal year 2015
18	and subsequent fiscal years, the fee amounts es-
19	tablished in paragraph (1) shall be adjusted by
20	the Secretary by notice, published in the Fed-
21	eral Register, for a fiscal year by the amount
22	equal to the sum of—
23	"(i) 1;
24	"(ii) the average annual percent
25	change in the cost, per full-time equivalent

1	position of the Food and Drug Administra-
2	tion, of all personnel compensation and
3	benefits paid with respect to such positions
4	for the first 3 years of the preceding 4 fis-
5	cal years, multiplied by the proportion of
6	personnel compensation and benefits costs
7	to total costs of an average full-time equiv-
8	alent position of the Food and Drug Ad-
9	ministration for the first 3 years of the
10	preceding 4 fiscal years; plus
11	"(iii) the average annual percent
12	change that occurred in the Consumer
13	Price Index for urban consumers (U.S.
14	City Average; Not Seasonally Adjusted; All
15	items; Annual Index) for the first 3 years
16	of the preceding 4 years of available data
17	multiplied by the proportion of all costs
18	other than personnel compensation and
19	benefits costs to total costs of an average
20	full-time equivalent position of the Food
21	and Drug Administration for the first 3
22	years of the preceding 4 fiscal years.
23	"(B) Compounded basis.—The adjust-
24	ment made each fiscal year under subparagraph
25	(A) shall be added on a compounded basis to

1	the sum of all adjustments made each fiscal
2	year after fiscal year 2014 under subparagraph
3	(A).
4	"(3) Small business adjustment factor.—
5	The small business adjustment factor described in
6	this paragraph shall be an amount established by
7	the Secretary for each fiscal year based on the Sec-
8	retary's estimate of—
9	"(A) the number of small businesses that
10	will pay a reduced establishment fee for such
11	fiscal year; and
12	"(B) the adjustment to the establishment
13	fee necessary to achieve total fees equaling the
14	total fees that the Secretary would have col-
15	lected if no entity qualified for the small busi-
16	ness exception in paragraph (4).
17	"(4) Exception for small businesses.—
18	"(A) In GENERAL.—In the case of an
19	outsourcing facility with gross annual sales of
20	\$1,000,000 or less in the 12 months ending
21	April 1 of the fiscal year immediately preceding
22	the fiscal year in which the fees under this sec-
23	tion are assessed, the amount of the establish-
24	ment fee under subsection (b) for a fiscal year

1	shall be equal to $1/3\$ of the amount calculated
2	under paragraph (1)(A)(i) for such fiscal year.
3	"(B) APPLICATION.—To qualify for the ex-
4	ception under this paragraph, a small business
5	shall submit to the Secretary a written request
6	for such exception, in a format specified by the
7	Secretary in guidance, certifying its gross an-
8	nual sales for the 12 months ending April 1 of
9	the fiscal year immediately preceding the fiscal
10	year in which fees under this subsection are as-
11	sessed. Any such application shall be submitted
12	to the Secretary not later than April 30 of such
13	immediately preceding fiscal year.
14	"(5) Crediting of Fees.—In establishing the
15	small business adjustment factor under paragraph
16	(3) for a fiscal year, the Secretary shall—
17	"(A) provide for the crediting of fees from
18	the previous year to the next year if the Sec-
19	retary overestimated the amount of the small
20	business adjustment factor for such previous
21	fiscal year; and
22	"(B) consider the need to account for any
23	adjustment of fees and such other factors as
24	the Secretary determines appropriate.

1	"(d) Use of Fees.—The Secretary shall make all
2	of the fees collected pursuant to subparagraphs (A) and
3	(B) of subsection (a)(1) available solely to pay for the
4	costs of oversight of outsourcing facilities.
5	"(e) Supplement Not Supplant.—Funds received
6	by the Secretary pursuant to this section shall be used
7	to supplement and not supplant any other Federal funds
8	available to carry out the activities described in this sec-
9	tion.
10	"(f) Crediting and Availability of Fees.—Fees
11	authorized under this section shall be collected and avail-
12	able for obligation only to the extent and in the amount
13	provided in advance in appropriations Acts. Such fees are
14	authorized to remain available until expended. Such sums
15	as may be necessary may be transferred from the Food
16	and Drug Administration salaries and expenses appropria-
17	tion account without fiscal year limitation to such appro-
18	priation account for salaries and expenses with such fiscal
19	year limitation. The sums transferred shall be available
20	solely for the purpose of paying the costs of oversight of
21	outsourcing facilities.
22	"(g) Collection of Fees.—
23	"(1) Establishment fee.—An outsourcing
24	facility shall remit the establishment fee due under
25	this section in a fiscal year when submitting a reg-

1	istration pursuant to section 503B(b) for such fiscal
2	year.
3	"(2) Reinspection fee.—The Secretary shall
4	specify in the Federal Register notice described in
5	subsection (b)(2) the manner in which reinspection
6	fees assessed under this section shall be collected
7	and the timeline for payment of such fees. Such a
8	fee shall be collected after the Secretary has con-
9	ducted a reinspection of the outsourcing facility in-
10	volved.
11	"(3) Effect of failure to pay fees.—
12	"(A) Registration.—An outsourcing fa-
13	cility shall not be considered registered under
14	section 503B(b) in a fiscal year until the date
15	that the outsourcing facility remits the estab-
16	lishment fee under this subsection for such fis-
17	cal year.
18	"(B) Misbranding.—All drugs manufac-
19	tured, prepared, propagated, compounded, or
20	processed by an outsourcing facility for which
21	any establishment fee or reinspection fee has
22	not been paid, as required by this section, shall
23	be deemed misbranded under section 502 until
24	the fees owed for such outsourcing facility
25	under this section have been paid.

1	"(4) Collection of unpaid fees.—In any
2	case where the Secretary does not receive payment
3	of a fee assessed under this section within 30 cal-
4	endar days after it is due, such fee shall be treated
5	as a claim of the United States Government subject
6	to provisions of subchapter II of chapter 37 of title
7	31, United States Code.
8	"(h) Annual Report to Congress.—Not later
9	than 120 calendar days after each fiscal year in which fees
10	are assessed and collected under this section, the Sec-
11	retary shall submit a report to the Committee on Health,
12	Education, Labor, and Pensions of the Senate and the
13	Committee on Energy and Commerce of the House of
14	Representatives, to include a description of fees assessed
15	and collected for such year, a summary description of enti-
16	ties paying the fees, a description of the hiring and place-
17	ment of new staff, a description of the use of fee resources
18	to support inspecting outsourcing facilities, and the num-
19	ber of inspections and reinspections of such facilities per-
20	formed each year.
21	"(i) Authorization of Appropriations.—For fis-
22	cal year 2014 and each subsequent fiscal year, there is
23	authorized to be appropriated for fees under this section
24	an amount equivalent to the total amount of fees assessed
25	for such fiscal year under this section.".

## 1 SEC. 103. PENALTIES.

- 2 (a) Prohibited Acts.—Section 301 (21 U.S.C.
- 3 331) is amended by adding at the end the following:
- 4 "(ccc)(1) The resale of a compounded drug that is
- 5 labeled 'not for resale' in accordance with section 503B.
- 6 "(2) With respect to a drug to be compounded pursu-
- 7 ant to section 503A or 503B, the intentional falsification
- 8 of a prescription, as applicable.
- 9 "(3) The failure to report drugs or adverse events
- 10 by an entity that is registered in accordance with sub-
- 11 section (b) of section 503B.".
- 12 (b) MISBRANDED DRUGS.—Section 502 (21 U.S.C.
- 13 352) is amended by adding at the end the following:
- 14 "(bb) If the advertising or promotion of a com-
- 15 pounded drug is false or misleading in any particular.".
- 16 SEC. 104. REGULATIONS.
- 17 In promulgating any regulations to implement this
- 18 title (and the amendments made by this title), the Sec-
- 19 retary of Health and Human Services shall—
- 20 (1) issue a notice of proposed rulemaking that
- 21 includes the proposed regulation;
- 22 (2) provide a period of not less than 60 cal-
- endar days for comments on the proposed regula-
- 24 tion; and
- 25 (3) publish the final regulation not more than
- 26 18 months following publication of the proposed rule

1	and not less than 30 calendar days before the effec-
2	tive date of such final regulation.
3	SEC. 105. ENHANCED COMMUNICATION.
4	(a) Submissions From State Boards of Phar-
5	MACY.—In a manner specified by the Secretary of Health
6	and Human Services (referred to in this section as the
7	"Secretary"), the Secretary shall receive submissions from
8	State boards of pharmacy—
9	(1) describing actions taken against
10	compounding pharmacies, as described in subsection
11	(b); or
12	(2) expressing concerns that a compounding
13	pharmacy may be acting contrary to section 503A of
14	the Federal Food, Drug, and Cosmetic Act (21
15	U.S.C. 353a).
16	(b) Content of Submissions From State
17	BOARDS OF PHARMACY.—An action referred to in sub-
18	section (a)(1) is, with respect to a pharmacy that com-
19	pounds drugs, any of the following:
20	(1) The issuance of a warning letter, or the im-
21	position of sanctions or penalties, by a State for vio-
22	lations of a State's pharmacy regulations pertaining
23	to compounding.
24	(2) The suspension or revocation of a State-
25	issued pharmacy license or registration for violations

1	of a State's pharmacy regulations pertaining to
2	compounding.
3	(3) The recall of a compounded drug due to
4	concerns relating to the quality or purity of such
5	drug.
6	(c) Consultation.—The Secretary shall implement
7	subsection (a) in consultation with the National Associa-
8	tion of Boards of Pharmacy.
9	(d) Notifying State Boards of Pharmacy.—The
10	Secretary shall immediately notify State boards of phar-
11	macy when—
12	(1) the Secretary receives a submission under
13	subsection $(a)(1)$ ; or
14	(2) the Secretary makes a determination that a
15	pharmacy is acting contrary to section 503A of the
16	Federal Food, Drug, and Cosmetic Act.
17	SEC. 106. SEVERABILITY.
18	(a) In General.—Section 503A (21 U.S.C. 353a)
19	is amended —
20	(1) in subsection (a), in the matter preceding
21	paragraph (1), by striking "unsolicited";
22	(2) by striking subsection (e);
23	(3) by redesignating subsections (d) through (f)
24	as subsections (c) through (e), respectively; and

1	(4) in subsection $(b)(1)(A)(i)(III)$ , by striking
2	"subsection (d)" and inserting "subsection (c)".
3	(b) SEVERABILITY.—If any provision of this Act (in-
4	cluding the amendments made by this Act) is declared un-
5	constitutional, or the applicability of this Act (including
6	the amendments made by this Act) to any person or cir-
7	cumstance is held invalid, the constitutionality of the re-
8	mainder of this Act (including the amendments made by
9	this Act) and the applicability thereof to other persons and
10	circumstances shall not be affected.
11	SEC. 107. GAO STUDY.
12	(a) STUDY.—Not later than 36 months after the date
13	of the enactment of this Act, the Comptroller General of
14	the United States shall submit to Congress a report on
15	pharmacy compounding and the adequacy of State and
16	Federal efforts to assure the safety of compounded drugs.
17	(b) CONTENTS.—The report required under this sec-
18	tion shall include—
19	(1) a review of pharmacy compounding in each
20	State, and the settings in which such compounding
21	occurs;
22	(2) a review of the State laws and policies gov-
23	erning pharmacy compounding, including enforce-
24	ment of State laws and policies;

1	(3) an assessment of the available tools to per-
2	mit purchasers of compounded drugs to determine
3	the safety and quality of such drugs;
4	(4) an evaluation of the effectiveness of the
5	communication among States and between States
6	and the Food and Drug Administration regarding
7	compounding; and
8	(5) an evaluation of the Food and Drug Admin-
9	istration's implementation of sections 503A and
10	503B of the Federal Food, Drug, and Cosmetic Act.
11	TITLE II—DRUG SUPPLY CHAIN
12	SECURITY
13	SEC. 201. SHORT TITLE.
14	This title may be cited as the "Drug Supply Chain
15	Security Act".
16	SEC. 202. PHARMACEUTICAL DISTRIBUTION SUPPLY
17	CHAIN.
18	Chapter V (21 U.S.C. 351 et seq.) is amended by
19	adding at the end the following:
20	"Subchapter H—Pharmaceutical Distribution
21	Supply Chain
22	"SEC. 581. DEFINITIONS.
23	"In this subchapter:

1	"(1) Affiliate.—The term 'affiliate' means a
2	business entity that has a relationship with a second
3	business entity if, directly or indirectly—
4	"(A) one business entity controls, or has
5	the power to control, the other business entity;
6	or
7	"(B) a third party controls, or has the
8	power to control, both of the business entities.
9	"(2) Authorized.—The term 'authorized'
10	means—
11	"(A) in the case of a manufacturer or re-
12	packager, having a valid registration in accord-
13	ance with section 510;
14	"(B) in the case of a wholesale distributor,
15	having a valid license under State law or sec-
16	tion 583, in accordance with section 582(a)(6),
17	and complying with the licensure reporting re-
18	quirements under section 503(e), as amended
19	by the Drug Supply Chain Security Act;
20	"(C) in the case of a third-party logistics
21	provider, having a valid license under State law
22	or section 584(a)(1), in accordance with section
23	582(a)(7), and complying with the licensure re-
24	porting requirements under section 584(b); and

1	"(D) in the case of a dispenser, having a
2	valid license under State law.
3	"(3) DISPENSER.—The term 'dispenser'—
4	"(A) means a retail pharmacy, hospital
5	pharmacy, a group of chain pharmacies under
6	common ownership and control that do not act
7	as a wholesale distributor, or any other person
8	authorized by law to dispense or administer
9	prescription drugs, and the affiliated ware-
10	houses or distribution centers of such entities
11	under common ownership and control that do
12	not act as a wholesale distributor; and
13	"(B) does not include a person who dis-
14	penses only products to be used in animals in
15	accordance with section 512(a)(5).
16	"(4) DISPOSITION.—The term 'disposition',
17	with respect to a product within the possession or
18	control of an entity, means the removal of such
19	product from the pharmaceutical distribution supply
20	chain, which may include disposal or return of the
21	product for disposal or other appropriate handling
22	and other actions, such as retaining a sample of the
23	product for further additional physical examination
24	or laboratory analysis of the product by a manufac-
25	turer or regulatory or law enforcement agency.

1	"(5) DISTRIBUTE OR DISTRIBUTION.—The
2	term 'distribute' or 'distribution' means the sale,
3	purchase, trade, delivery, handling, storage, or re-
4	ceipt of a product, and does not include the dis-
5	pensing of a product pursuant to a prescription exe-
6	cuted in accordance with section $503(b)(1)$ or the
7	dispensing of a product approved under section
8	512(b).
9	"(6) Exclusive distributor.—The term 'ex-
10	clusive distributor' means the wholesale distributor
11	that directly purchased the product from the manu-
12	facturer and is the sole distributor of that manufac-
13	turer's product to a subsequent repackager, whole-
14	sale distributor, or dispenser.
15	"(7) Homogeneous case.—The term 'homo-
16	geneous case' means a sealed case containing only
17	product that has a single National Drug Code num-
18	ber belonging to a single lot.
19	"(8) Illegitimate product.—The term 'ille-
20	gitimate product' means a product for which credible
21	evidence shows that the product—
22	"(A) is counterfeit, diverted, or stolen;
23	"(B) is intentionally adulterated such that
24	the product would result in serious adverse
25	health consequences or death to humans;

1	"(C) is the subject of a fraudulent trans-
2	action; or
3	"(D) appears otherwise unfit for distribu-
4	tion such that the product would be reasonably
5	likely to result in serious adverse health con-
6	sequences or death to humans.
7	"(9) Licensed.—The term 'licensed' means—
8	"(A) in the case of a wholesale distributor,
9	having a valid license in accordance with section
10	503(e) or section 582(a)(6), as applicable;
11	"(B) in the case of a third-party logistics
12	provider, having a valid license in accordance
13	with section 584(a) or section 582(a)(7), as ap-
14	plicable; and
15	"(C) in the case of a dispenser, having a
16	valid license under State law.
17	"(10) Manufacturer.—The term 'manufac-
18	turer' means, with respect to a product—
19	"(A) a person that holds an application ap-
20	proved under section 505 or a license issued
21	under section 351 of the Public Health Service
22	Act for such product, or if such product is not
23	the subject of an approved application or li-
24	cense, the person who manufactured the prod-
25	uct;

1	"(B) a co-licensed partner of the person
2	described in subparagraph (A) that obtains the
3	product directly from a person described in this
4	subparagraph or subparagraph (A) or (C); or
5	"(C) an affiliate of a person described in
6	subparagraph (A) or (B) that receives the prod-
7	uct directly from a person described in this sub-
8	paragraph or subparagraph (A) or (B).
9	"(11) Package.—
10	"(A) IN GENERAL.—The term 'package'
11	means the smallest individual saleable unit of
12	product for distribution by a manufacturer or
13	repackager that is intended by the manufac-
14	turer for ultimate sale to the dispenser of such
15	product.
16	"(B) Individual saleable unit.—For
17	purposes of this paragraph, an 'individual sale-
18	able unit' is the smallest container of product
19	introduced into commerce by the manufacturer
20	or repackager that is intended by the manufac-
21	turer or repackager for individual sale to a dis-
22	penser.
23	"(12) Prescription drug.—The term 'pre-
24	scription drug' means a drug for human use subject
25	to section 503(b)(1).

1	"(13) Product.—The term 'product' means a
2	prescription drug in a finished dosage form for ad-
3	ministration to a patient without substantial further
4	manufacturing (such as capsules, tablets, and
5	lyophilized products before reconstitution), but for
6	purposes of section 582, does not include blood or
7	blood components intended for transfusion, radio-
8	active drugs or radioactive biological products (as
9	defined in section 600.3(ee) of title 21, Code of Fed-
10	eral Regulations) that are regulated by the Nuclear
11	Regulatory Commission or by a State pursuant to
12	an agreement with such Commission under section
13	274 of the Atomic Energy Act of 1954 (42 U.S.C.
14	2021), imaging drugs, an intravenous product de-
15	scribed in clause (xiv), (xv), or (xvi) of paragraph
16	(24)(B), any medical gas (as defined in section 575),
17	homeopathic drugs marketed in accordance with ap-
18	plicable guidance under this Act, or a drug com-
19	pounded in compliance with section 503A or 503B.
20	"(14) Product identifier.—The term 'prod-
21	uct identifier' means a standardized graphic that in-
22	cludes, in both human-readable form and on a ma-
23	chine-readable data carrier that conforms to the
24	standards developed by a widely recognized inter-
25	national standards development organization, the

1	standardized numerical identifier, lot number, and
2	expiration date of the product.
3	"(15) QUARANTINE.—The term 'quarantine'
4	means the storage or identification of a product, to
5	prevent distribution or transfer of the product, in a
6	physically separate area clearly identified for such
7	use or through other procedures.
8	"(16) Repackager.—The term 'repackager'
9	means a person who owns or operates an establish-
10	ment that repacks and relabels a product or package
11	for—
12	"(A) further sale; or
13	"(B) distribution without a further trans-
14	action.
15	"(17) Return.—The term 'return' means pro-
16	viding product to the authorized immediate trading
17	partner from which such product was purchased or
18	received, or to a returns processor or reverse logis-
19	tics provider for handling of such product.
20	"(18) Returns processor or reverse lo-
21	GISTICS PROVIDER.—The term 'returns processor' or
22	'reverse logistics provider' means a person who owns
23	or operates an establishment that dispositions or
24	otherwise processes saleable or nonsaleable product
25	received from an authorized trading partner such

1	that the product may be processed for credit to the
2	purchaser, manufacturer, or seller or disposed of for
3	no further distribution.
4	"(19) Specific patient need.—The term
5	'specific patient need' refers to the transfer of a
6	product from one pharmacy to another to fill a pre-
7	scription for an identified patient. Such term does
8	not include the transfer of a product from one phar-
9	macy to another for the purpose of increasing or re-
10	plenishing stock in anticipation of a potential need
11	"(20) Standardized numerical identi-
12	FIER.—The term 'standardized numerical identifier
13	means a set of numbers or characters used to
14	uniquely identify each package or homogenous case
15	that is composed of the National Drug Code that
16	corresponds to the specific product (including the
17	particular package configuration) combined with a
18	unique alphanumeric serial number of up to 20
19	characters.
20	"(21) Suspect Product.—The term 'suspect
21	product' means a product for which there is reason
22	to believe that such product—
23	"(A) is potentially counterfeit, diverted, or
24	stolen;

1	"(B) is potentially intentionally adulterated
2	such that the product would result in serious
3	adverse health consequences or death to hu-
4	mans;
5	"(C) is potentially the subject of a fraudu-
6	lent transaction; or
7	"(D) appears otherwise unfit for distribu-
8	tion such that the product would result in seri-
9	ous adverse health consequences or death to hu-
10	mans.
11	"(22) Third-party logistics provider.—
12	The term 'third-party logistics provider' means an
13	entity that provides or coordinates warehousing, or
14	other logistics services of a product in interstate
15	commerce on behalf of a manufacturer, wholesale
16	distributor, or dispenser of a product, but does not
17	take ownership of the product, nor have responsi-
18	bility to direct the sale or disposition of the product.
19	"(23) Trading Partner.—The term 'trading
20	partner' means—
21	"(A) a manufacturer, repackager, whole-
22	sale distributor, or dispenser from whom a
23	manufacturer, repackager, wholesale dis-
24	tributor, or dispenser accepts direct ownership
25	of a product or to whom a manufacturer, re-

1	packager, wholesale distributor, or dispenser
2	transfers direct ownership of a product; or
3	"(B) a third-party logistics provider from
4	whom a manufacturer, repackager, wholesale
5	distributor, or dispenser accepts direct posses-
6	sion of a product or to whom a manufacturer,
7	repackager, wholesale distributor, or dispenser
8	transfers direct possession of a product.
9	"(24) Transaction.—
10	"(A) IN GENERAL.—The term 'transaction'
11	means the transfer of product between persons
12	in which a change of ownership occurs.
13	"(B) Exemptions.—The term 'trans-
14	action' does not include—
15	"(i) intracompany distribution of any
16	product between members of an affiliate or
17	within a manufacturer;
18	"(ii) the distribution of a product
19	among hospitals or other health care enti-
20	ties that are under common control;
21	"(iii) the distribution of a product for
22	emergency medical reasons including a
23	public health emergency declaration pursu-
24	ant to section 319 of the Public Health
25	Service Act. except that a drug shortage

1	not caused by a public health emergency
2	shall not constitute an emergency medical
3	reason;
4	"(iv) the dispensing of a product pur-
5	suant to a prescription executed in accord-
6	ance with section 503(b)(1);
7	"(v) the distribution of product sam-
8	ples by a manufacturer or a licensed
9	wholesale distributor in accordance with
10	section 503(d);
11	"(vi) the distribution of blood or blood
12	components intended for transfusion;
13	"(vii) the distribution of minimal
14	quantities of product by a licensed retail
15	pharmacy to a licensed practitioner for of-
16	fice use;
17	"(viii) the sale, purchase, or trade of
18	a drug or an offer to sell, purchase, or
19	trade a drug by a charitable organization
20	described in section $501(c)(3)$ of the Inter-
21	nal Revenue Code of 1986 to a nonprofit
22	affiliate of the organization to the extent
23	otherwise permitted by law;
24	"(ix) the distribution of a product
25	pursuant to the sale or merger of a phar-

1	macy or pharmacies or a wholesale dis-
2	tributor or wholesale distributors, except
3	that any records required to be maintained
4	for the product shall be transferred to the
5	new owner of the pharmacy or pharmacies
6	or wholesale distributor or wholesale dis-
7	tributors;
8	"(x) the dispensing of a product ap-
9	proved under section 512(c);
10	"(xi) products transferred to or from
11	any facility that is licensed by the Nuclear
12	Regulatory Commission or by a State pur-
13	suant to an agreement with such Commis-
14	sion under section 274 of the Atomic En-
15	ergy Act of 1954 (42 U.S.C. 2021);
16	"(xii) a combination product that is
17	not subject to approval under section 505
18	or licensure under section 351 of the Pub-
19	lic Health Service Act, and that is—
20	"(I) a product comprised of a de-
21	vice and 1 or more other regulated
22	components (such as a drug/device,
23	biologic/device, or drug/device/biologic)
24	that are physically, chemically, or oth-

1	erwise combined or mixed and pro-
2	duced as a single entity;
3	"(II) 2 or more separate prod-
4	ucts packaged together in a single
5	package or as a unit and comprised of
6	a drug and device or device and bio-
7	logical product; or
8	"(III) 2 or more finished medical
9	devices plus one or more drug or bio-
10	logical products that are packaged to-
11	gether in what is referred to as a
12	'medical convenience kit' as described
13	in clause (xiii);
14	"(xiii) the distribution of a collection
15	of finished medical devices, which may in-
16	clude a product or biological product, as-
17	sembled in kit form strictly for the conven-
18	ience of the purchaser or user (referred to
19	in this clause as a 'medical convenience
20	kit') if—
21	"(I) the medical convenience kit
22	is assembled in an establishment that
23	is registered with the Food and Drug
24	Administration as a device manufac-

1	turer in accordance with section
2	510(b)(2);
3	"(II) the medical convenience kit
4	does not contain a controlled sub-
5	stance that appears in a schedule con-
6	tained in the Comprehensive Drug
7	Abuse Prevention and Control Act of
8	1970;
9	"(III) in the case of a medical
10	convenience kit that includes a prod-
11	uct, the person that manufacturers
12	the kit—
13	"(aa) purchased such prod-
13 14	"(aa) purchased such prod- uct directly from the pharma-
14	uct directly from the pharma-
14 15	uct directly from the pharma- ceutical manufacturer or from a
<ul><li>14</li><li>15</li><li>16</li></ul>	uct directly from the pharma- ceutical manufacturer or from a wholesale distributor that pur-
<ul><li>14</li><li>15</li><li>16</li><li>17</li></ul>	uct directly from the pharma- ceutical manufacturer or from a wholesale distributor that pur- chased the product directly from
<ul><li>14</li><li>15</li><li>16</li><li>17</li><li>18</li></ul>	uct directly from the pharmaceutical manufacturer or from a wholesale distributor that purchased the product directly from the pharmaceutical manufac-
<ul><li>14</li><li>15</li><li>16</li><li>17</li><li>18</li><li>19</li></ul>	uct directly from the pharmaceutical manufacturer or from a wholesale distributor that purchased the product directly from the pharmaceutical manufacturer; and
14 15 16 17 18 19 20	uct directly from the pharmaceutical manufacturer or from a wholesale distributor that purchased the product directly from the pharmaceutical manufacturer; and "(bb) does not alter the pri-
14 15 16 17 18 19 20 21	uct directly from the pharmaceutical manufacturer or from a wholesale distributor that purchased the product directly from the pharmaceutical manufacturer; and  "(bb) does not alter the primary container or label of the

1	"(IV) in the case of a medical
2	convenience kit that includes a prod-
3	uct, the product is—
4	"(aa) an intravenous solu-
5	tion intended for the replenish-
6	ment of fluids and electrolytes;
7	"(bb) a product intended to
8	maintain the equilibrium of water
9	and minerals in the body;
10	"(cc) a product intended for
11	irrigation or reconstitution;
12	"(dd) an anesthetic;
13	"(ee) an anticoagulant;
14	"(ff) a vasopressor; or
15	"(gg) a sympathomimetic;
16	"(xiv) the distribution of an intra-
17	venous product that, by its formulation, is
18	intended for the replenishment of fluids
19	and electrolytes (such as sodium, chloride,
20	and potassium) or calories (such as dex-
21	trose and amino acids);
22	"(xv) the distribution of an intra-
23	venous product used to maintain the equi-
24	librium of water and minerals in the body,
25	such as dialysis solutions;

1	"(xvi) the distribution of a product
2	that is intended for irrigation, or sterile
3	water, whether intended for such purposes
4	or for injection;
5	"(xvii) the distribution of a medical
6	gas (as defined in section 575); or
7	"(xviii) the distribution or sale of any
8	licensed product under section 351 of the
9	Public Health Service Act that meets the
10	definition of a device under section 201(h).
11	"(25) Transaction History.—The term
12	'transaction history' means a statement in paper or
13	electronic form, including the transaction informa-
14	tion for each prior transaction going back to the
15	manufacturer of the product.
16	"(26) Transaction information.—The term
17	'transaction information' means—
18	"(A) the proprietary or established name
19	or names of the product;
20	"(B) the strength and dosage form of the
21	product;
22	"(C) the National Drug Code number of
23	the product;
24	"(D) the container size;
25	"(E) the number of containers;

1	"(F) the lot number of the product;
2	"(G) the date of the transaction;
3	"(H) the date of the shipment, if more
4	than 24 hours after the date of the transaction
5	"(I) the business name and address of the
6	person from whom ownership is being trans-
7	ferred; and
8	"(J) the business name and address of the
9	person to whom ownership is being transferred
10	"(27) Transaction statement.—The 'trans-
11	action statement' is a statement, in paper or elec-
12	tronic form, that the entity transferring ownership
13	in a transaction—
14	"(A) is authorized as required under the
15	Drug Supply Chain Security Act;
16	"(B) received the product from a person
17	that is authorized as required under the Drug
18	Supply Chain Security Act;
19	"(C) received transaction information and
20	a transaction statement from the prior owner of
21	the product, as required under section 582;
22	"(D) did not knowingly ship a suspect or
23	illegitimate product;

1	"(E) had systems and processes in place to
2	comply with verification requirements under
3	section 582;
4	"(F) did not knowingly provide false trans-
5	action information; and
6	"(G) did not knowingly alter the trans-
7	action history.
8	"(28) Verification or verify.—The term
9	'verification' or 'verify' means determining whether
10	the product identifier affixed to, or imprinted upon,
11	a package or homogeneous case corresponds to the
12	standardized numerical identifier or lot number and
13	expiration date assigned to the product by the man-
14	ufacturer or the repackager, as applicable in accord-
15	ance with section 582.
16	"(29) Wholesale distributor.—The term
17	'wholesale distributor' means a person (other than a
18	manufacturer, a manufacturer's co-licensed partner,
19	a third-party logistics provider, or repackager) en-
20	gaged in wholesale distribution (as defined in section
21	503(e)(4), as amended by the Drug Supply Chain
22	Security Act).
23	"SEC. 582. REQUIREMENTS.
24	"(a) In General.—

"(1) OTHER ACTIVITIES.—Each manufacturer, repackager, wholesale distributor, and dispenser shall comply with the requirements set forth in this section with respect to the role of such manufacturer, repackager, wholesale distributor, or dispenser in a transaction involving product. If an entity meets the definition of more than one of the entities listed in the preceding sentence, such entity shall comply with all applicable requirements in this section, but shall not be required to duplicate requirements.

## "(2) Initial standards.—

"(A) IN GENERAL.—The Secretary shall, in consultation with other appropriate Federal officials, manufacturers, repackagers, wholesale distributors, dispensers, and other pharmaceutical distribution supply chain stakeholders, issue a draft guidance document that establishes standards for the interoperable exchange of transaction information, transaction history, and transaction statements, in paper or electronic format, for compliance with this subsection and subsections (b), (c), (d), and (e). In establishing such standards, the Secretary shall consider the feasibility of establishing standardized documentation to be used by members of

1	the pharmaceutical distribution supply chain to
2	convey the transaction information, transaction
3	history, and transaction statement to the subse-
4	quent purchaser of a product and to facilitate
5	the exchange of lot level data. The standards
6	established under this paragraph shall take into
7	consideration the standards established under
8	section 505D and shall comply with a form and
9	format developed by a widely recognized inter-
10	national standards development organization.
11	"(B) Public input.—Prior to issuing the
12	draft guidance under subparagraph (A), the
13	Secretary shall gather comments and informa-
14	tion from stakeholders and maintain such com-
15	ments and information in a public docket for at
16	least 60 days prior to issuing such guidance.
17	"(C) Publication.—The Secretary shall
18	publish the standards established under sub-
19	paragraph (A) not later than 1 year after the
20	date of enactment of the Drug Supply Chain
21	Security Act.
22	"(3) Waivers, exceptions, and exemp-
23	TIONS.—
24	"(A) IN GENERAL.—Not later than 2 years
25	after the date of enactment of the Drug Supply

1	Chain Security Act, the Secretary shall, by
2	guidance—
3	"(i) establish a process by which an
4	authorized manufacturer, repackager,
5	wholesale distributor, or dispenser may re-
6	quest a waiver from any of the require-
7	ments set forth in this section, which the
8	Secretary may grant if the Secretary deter-
9	mines that such requirements would result
10	in an undue economic hardship or for
11	emergency medical reasons, including a
12	public health emergency declaration pursu-
13	ant to section 319 of the Public Health
14	Service Act;
15	"(ii) establish a process by which the
16	Secretary determines exceptions, and a
17	process through which a manufacturer or
18	repackager may request such an exception,
19	to the requirements relating to product
20	identifiers if a product is packaged in a
21	container too small or otherwise unable to
22	accommodate a label with sufficient space
23	to bear the information required for com-
24	pliance with this section; and

1	"(iii) establish a process by which the
2	Secretary may determine other products or
3	transactions that shall be exempt from the
4	requirements of this section.
5	"(B) Content.—The guidance issued
6	under subparagraph (A) shall include a process
7	for the biennial review and renewal of such
8	waivers, exceptions, and exemptions, as applica-
9	ble.
10	"(C) Process.—In issuing the guidance
11	under this paragraph, the Secretary shall pro-
12	vide an effective date that is not later than 180
13	days prior to the date on which manufacturers
14	are required to affix or imprint a product iden-
15	tifier to each package and homogenous case of
16	product intended to be introduced in a trans-
17	action into commerce consistent with this sec-
18	tion.
19	"(4) Self-executing requirements.—Ex-
20	cept where otherwise specified, the requirements of
21	this section may be enforced without further regula-
22	tions or guidance from the Secretary.
23	"(5) Grandfathering product.—
24	"(A) Product identifier.—Not later
25	than 2 years after the date of enactment of the

1	Drug Supply Chain Security Act, the Secretary
2	shall finalize guidance specifying whether and
3	under what circumstances product that is not
4	labeled with a product identifier and that is in
5	the pharmaceutical distribution supply chain at
6	the time of the effective date of the require-
7	ments of this section shall be exempted from
8	the requirements of this section.
9	"(B) Tracing.—For a product that en-
10	tered the pharmaceutical distribution supply
11	chain prior to January 1, 2015—
12	"(i) authorized trading partners shall
13	be exempt from providing transaction in-
14	formation as required under subsections
15	(b)(1)(A)(i), (c)(1)(A)(ii), (d)(1)(A)(ii),
16	and (e)(1)(A)(ii);
17	"(ii) transaction history required
18	under this section shall begin with the
19	owner of such product on such date; and
20	"(iii) the owners of such product on
21	such date shall be exempt from asserting
22	receipt of transaction information and
23	transaction statement from the prior owner
24	as required under this section.

1	"(6) Wholesale distributor licenses.—
2	Notwithstanding section 581(9)(A), until the effec-
3	tive date of the wholesale distributor licensing regu-
4	lations under section 583, the term 'licensed' or 'au-
5	thorized', as it relates to a wholesale distributor with
6	respect to prescription drugs, shall mean a wholesale
7	distributor with a valid license under State law.
8	"(7) Third-party logistics provider li-
9	CENSES.—Until the effective date of the third-party
10	logistics provider licensing regulations under section
11	584, a third-party logistics provider shall be consid-
12	ered 'licensed' under section 581(9)(B) unless the
13	Secretary has made a finding that the third-party lo-
14	gistics provider does not utilize good handling and
15	distribution practices and publishes notice thereof.
16	"(8) Label Changes.—Changes made to pack-
17	age labels solely to incorporate the product identifier
18	may be submitted to the Secretary in the annual re-
19	port of an establishment, in accordance with section
20	314.70(d) of chapter 21, Code of Federal Regula-
21	tions (or any successor regulation).
22	"(9) Product identifiers.—With respect to
23	any requirement relating to product identifiers under
24	this subchapter—

1	"(A) unless the Secretary allows, through
2	guidance, the use of other technologies for data
3	instead of or in addition to the technologies de-
4	scribed in clauses (i) and (ii), the applicable
5	data—
6	"(i) shall be included in a 2-dimen-
7	sional data matrix barcode when affixed to,
8	or imprinted upon, a package; and
9	"(ii) shall be included in a linear or 2-
10	dimensional data matrix barcode when af-
11	fixed to, or imprinted upon, a homo-
12	geneous case; and
13	"(B) verification of the product identifier
14	may occur by using human-readable or ma-
15	chine-readable methods.
16	"(b) Manufacturer Requirements.—
17	"(1) Product tracing.—
18	"(A) In General.—Beginning not later
19	than January 1, 2015, a manufacturer shall—
20	"(i) prior to, or at the time of, each
21	transaction in which such manufacturer
22	transfers ownership of a product, provide
23	the subsequent owner with transaction his-
24	tory, transaction information, and a trans-

1	action statement, in a single document in
2	an paper or electronic format; and
3	"(ii) capture the transaction informa-
4	tion (including lot level information),
5	transaction history, and transaction state-
6	ment for each transaction and maintain
7	such information, history, and statement
8	for not less than 6 years after the date of
9	the transaction.
10	"(B) REQUESTS FOR INFORMATION.—
11	Upon a request by the Secretary or other ap-
12	propriate Federal or State official, in the event
13	of a recall or for the purpose of investigating a
14	suspect product or an illegitimate product, a
15	manufacturer shall, not later than 1 business
16	day, and not to exceed 48 hours, after receiving
17	the request, or in other such reasonable time as
18	determined by the Secretary, based on the cir-
19	cumstances of the request, provide the applica-
20	ble transaction information, transaction history,
21	and transaction statement for the product.
22	"(C) ELECTRONIC FORMAT.—
23	"(i) In General.—Beginning not
24	later than 4 years after the date of enact-
25	ment of the Drug Supply Chain Security

1	Act, except as provided under clause (ii), a
2	manufacturer shall provide the transaction
3	information, transaction history, and
4	transaction statement required under sub-
5	paragraph (A)(i) in electronic format.
6	"(ii) Exception.—A manufacturer
7	may continue to provide the transaction in-
8	formation, transaction history, and trans-
9	action statement required under subpara-
10	graph (A)(i) in a paper format to a li-
11	censed health care practitioner authorized
12	to prescribe medication under State law or
13	other licensed individual under the super-
14	vision or direction of such a practitioner
15	who dispenses product in the usual course
16	of professional practice.
17	"(2) Product identifier.—
18	"(A) IN GENERAL.—Beginning not later
19	than 4 years after the date of enactment of the
20	Drug Supply Chain Security Act, a manufac-
21	turer shall affix or imprint a product identifier
22	to each package and homogenous case of a
23	product intended to be introduced in a trans-
24	action into commerce. Such manufacturer shall

maintain the product identifier information for

25

1	such product for not less than 6 years after the
2	date of the transaction.
3	"(B) Exception.—A package that is re-
4	quired to have a standardized numerical identi-
5	fier is not required to have a unique device
6	identifier.
7	"(3) Authorized trading partners.—Be-
8	ginning not later than January 1, 2015, the trading
9	partners of a manufacturer may be only authorized
10	trading partners.
11	"(4) Verification.—Beginning not later than
12	January 1, 2015, a manufacturer shall have systems
13	in place to enable the manufacturer to comply with
14	the following requirements:
15	"(A) Suspect product.—
16	"(i) In General.—Upon making a
17	determination that a product in the posses-
18	sion or control of the manufacturer is a
19	suspect product, or upon receiving a re-
20	quest for verification from the Secretary
21	that has made a determination that a
22	product within the possession or control of
23	a manufacturer is a suspect product, a
24	manufacturer shall—

1	"(I) quarantine such product
2	within the possession or control of the
3	manufacturer from product intended
4	for distribution until such product is
5	cleared or dispositioned; and
6	"(II) promptly conduct an inves-
7	tigation in coordination with trading
8	partners, as applicable, to determine
9	whether the product is an illegitimate
10	product, which shall include validating
11	any applicable transaction history and
12	transaction information in the posses-
13	sion of the manufacturer and other-
14	wise investigating to determine wheth-
15	er the product is an illegitimate prod-
16	uct, and, beginning 4 years after the
17	date of enactment of the Drug Supply
18	Chain Security Act, verifying the
19	product at the package level, including
20	the standardized numerical identifier.
21	"(ii) CLEARED PRODUCT.—If the
22	manufacturer makes the determination
23	that a suspect product is not an illegit-
24	imate product, the manufacturer shall
25	promptly notify the Secretary, if applica-

1	ble, of such determination and such prod-
2	uct may be further distributed.
3	"(iii) Records.—A manufacturer
4	shall keep records of the investigation of a
5	suspect product for not less than 6 years
6	after the conclusion of the investigation.
7	"(B) Illegitimate product.—
8	"(i) In General.—Upon determining
9	that a product in the possession or control
10	of a manufacturer is an illegitimate prod-
11	uct, the manufacturer shall, in a manner
12	consistent with the systems and processes
13	of such manufacturer—
14	"(I) quarantine such product
15	within the possession or control of the
16	manufacturer from product intended
17	for distribution until such product is
18	dispositioned;
19	"(II) disposition the illegitimate
20	product within the possession or con-
21	trol of the manufacturer;
22	"(III) take reasonable and appro-
23	priate steps to assist a trading part-

1	uct not in the possession or control of
2	the manufacturer; and
3	"(IV) retain a sample of the
4	product for further physical examina-
5	tion or laboratory analysis of the
6	product by the manufacturer or Sec-
7	retary (or other appropriate Federal
8	or State official) upon request by the
9	Secretary (or other appropriate Fed-
10	eral or State official), as necessary
11	and appropriate.
12	"(ii) Making a notification.—
13	"(I) ILLEGITIMATE PRODUCT.—
14	Upon determining that a product in
15	the possession or control of the manu-
16	facturer is an illegitimate product, the
17	manufacturer shall notify the Sec-
18	retary and all immediate trading part-
19	ners that the manufacturer has reason
20	to believe may have received such ille-
21	gitimate product of such determina-
22	tion not later than 24 hours after
23	making such determination.
24	"(II) High risk of illegit-
25	IMACY.—A manufacturer shall notify

1	the Secretary and immediate trading
2	partners that the manufacturer has
3	reason to believe may have in the
4	trading partner's possession a product
5	manufactured by, or purported to be a
6	product manufactured by, the manu-
7	facturer not later than 24 hours after
8	determining or being notified by the
9	Secretary or a trading partner that
10	there is a high risk that such product
11	is an illegitimate product. For pur-
12	poses of this subclause, a 'high risk'
13	may include a specific high risk that
14	could increase the likelihood that ille-
15	gitimate product will enter the phar-
16	maceutical distribution supply chain
17	and other high risks as determined by
18	the Secretary in guidance pursuant to
19	subsection (h).
20	"(iii) Responding to a notifica-
21	TION.—Upon the receipt of a notification
22	from the Secretary or a trading partner
23	that a determination has been made that a
24	product is an illegitimate product, a manu-
25	facturer shall identify all illegitimate prod-

1	uct subject to such notification that is in
2	the possession or control of the manufac-
3	turer, including any product that is subse-
4	quently received, and shall perform the ac-
5	tivities described in subparagraph (A).
6	"(iv) TERMINATING A NOTIFICA-
7	TION.—Upon making a determination, in
8	consultation with the Secretary, that a no-
9	tification is no longer necessary, a manu-
10	facturer shall promptly notify immediate
11	trading partners that the manufacturer no-
12	tified pursuant to clause (ii) that such no-
13	tification has been terminated.
14	"(v) Records.—A manufacturer shall
15	keep records of the disposition of an illegit-
16	imate product for not less than 6 years
17	after the conclusion of the disposition.
18	"(C) Requests for verification.—Be-
19	ginning 4 years after the date of enactment of
20	the Drug Supply Chain Security Act, upon re-
21	ceiving a request for verification from an au-
22	thorized repackager, wholesale distributor, or
23	dispenser that is in possession or control of a
24	product such person believes to be manufac-
25	tured by such manufacturer, a manufacturer

shall, not later than 24 hours after receiving 1 2 the request for verification or in other such reasonable time as determined by the Secretary, 3 based on the circumstances of the request, notify the person making the request whether the 6 product identifier, including the standardized 7 numerical identifier, that is the subject of the 8 request corresponds to the product identifier af-9 fixed or imprinted by the manufacturer. If a 10 manufacturer responding to a request for 11 verification identifies a product identifier that 12 does not correspond to that affixed or imprinted 13 by the manufacturer, the manufacturer shall 14 treat such product as suspect product and con-15 duct an investigation as described in subpara-16 graph (A). If the manufacturer has reason to 17 believe the product is an illegitimate product, 18 the manufacturer shall advise the person mak-19 ing the request of such belief at the time such 20 manufacturer responds to the request for 21 verification. 22 "(D) ELECTRONIC DATABASE.—A manu-23 facturer may satisfy the requirements of this 24 paragraph by developing a secure electronic 25 database or utilizing a secure electronic database developed or operated by another entity. The owner of such database shall establish the requirements and processes to respond to requests and may provide for data access to other members of the pharmaceutical distribution supply chain, as appropriate. The development and operation of such a database shall not relieve a manufacturer of the requirement under this paragraph to respond to a request for verification submitted by means other than a secure electronic database.

"(E) SALEABLE RETURNED PRODUCT.—
Beginning 4 years after the date of enactment of the Drug Supply Chain Security Act (except as provided pursuant to subsection (a)(5)), upon receipt of a returned product that the manufacturer intends to further distribute, before further distributing such product, the manufacturer shall verify the product identifier, including the standardized numerical identifier, for each sealed homogeneous case of such product or, if such product is not in a sealed homogeneous case, verify the product identifier, including the standardized numerical identifier, on each package.

1	"(F) Nonsaleable returned prod-
2	UCT.—A manufacturer may return a nonsale-
3	able product to the manufacturer or repack-
4	ager, to the wholesale distributor from whom
5	such product was purchased, or to a person act-
6	ing on behalf of such a person, including a re-
7	turns processor, without providing the informa-
8	tion described in paragraph (1)(A)(i).
9	"(c) Wholesale Distributor Requirements.—
10	"(1) Product tracing.—
11	"(A) In General.—Beginning not later
12	than January 1, 2015, the following require-
13	ments shall apply to wholesale distributors:
14	"(i) A wholesale distributor shall not
15	accept ownership of a product unless the
16	previous owner prior to, or at the time of,
17	the transaction provides the transaction
18	history, transaction information, and a
19	transaction statement for the product, as
20	applicable under this subparagraph.
21	"(ii)(I)(aa) If the wholesale dis-
22	tributor purchased a product directly from
23	the manufacturer, the exclusive distributor
24	of the manufacturer, or a repackager that
25	purchased directly from the manufacturer,

1	then prior to, or at the time of, each trans-
2	action in which the wholesale distributor
3	transfers ownership of a product, the
4	wholesale distributor shall provide to the
5	subsequent purchaser—
6	"(AA) a transaction statement,
7	which shall state that such wholesale
8	distributor, or a member of the affil-
9	iate of such wholesale distributor, pur-
10	chased the product directly from the
11	manufacturer, exclusive distributor of
12	the manufacturer, or repackager that
13	purchased the product directly from
14	the manufacturer; and
15	"(BB) subject to subclause (II),
16	the transaction history and trans-
17	action information.
18	"(bb) The wholesale distributor shall
19	provide the transaction history, transaction
20	information, and transaction statement
21	under item (aa)—
22	"(AA) if provided to a dis-
23	penser, on a single document in a
24	paper or electronic format; and

1 "(BB) if pro	ovided to a
2 wholesale distribut	tor, through
3 any combination of	self-generated
4 paper, electronic da	ıta, or manu-
5 facturer-provided in	nformation on
6 the product package	<u>,</u>
7 "(II) For purposes of tra	ansactions de-
8 scribed in subclause (I), tra	ansaction his-
9 tory and transaction informa	tion shall not
be required to include the le	ot number of
11 the product, the initial transa	action date, or
the initial shipment date fro	m the manu-
facturer (as defined in subpa	ragraphs (F),
14 (G), and (H) of section 581(2	(6)).
15 "(iii) If the wholesale d	istributor did
not purchase a product dire	ctly from the
manufacturer, the exclusive	distributor of
the manufacturer, or a rep	oackager that
purchased directly from the 1	manufacturer,
as described in clause (ii), the	en prior to, or
21 at the time of, each transact	tion or subse-
quent transaction, the wholesa	ale distributor
shall provide to the subsequer	nt purchaser a
24 transaction statement, transa	action history,
and transaction information,	in a paper or

1	electronic format that complies with the
2	guidance document issued under sub-
3	section (a)(2).
4	"(iv) For the purposes of clause (iii),
5	the transaction history supplied shall begin
6	only with the wholesale distributor de-
7	scribed in clause (ii)(I), but the wholesale
8	distributor described in clause (iii) shall in-
9	form the subsequent purchaser that such
10	wholesale distributor received a direct pur-
11	chase statement from a wholesale dis-
12	tributor described in clause (ii)(I).
13	"(v) A wholesale distributor shall—
14	"(I) capture the transaction in-
15	formation (including lot level informa-
16	tion) consistent with the requirements
17	of this section, transaction history,
18	and transaction statement for each
19	transaction described in clauses (i),
20	(ii), and (iii) and maintain such infor-
21	mation, history, and statement for not
22	less than 6 years after the date of the
23	transaction; and
24	"(II) maintain the confidentiality
25	of the transaction information (includ-

1	ing any lot level information con-
2	sistent with the requirements of this
3	section), transaction history, and
4	transaction statement for a product in
5	a manner that prohibits disclosure to
6	any person other than the Secretary
7	or other appropriate Federal or State
8	official, except to comply with clauses
9	(ii) and (iii), and, as applicable, pur-
10	suant to an agreement under subpara-
11	graph (D).
12	"(B) Returns.—
13	"(i) Saleable returns.—Notwith-
14	standing subparagraph (A)(i), the fol-
15	lowing shall apply:
16	"(I) REQUIREMENTS.—Until the
17	date that is 6 years after the date of
18	enactment of the Drug Supply Chain
19	Security Act (except as provided pur-
20	suant to subsection (a)(5)), a whole-
21	sale distributor may accept returned
22	product from a dispenser or repack-
23	ager pursuant to the terms and condi-
24	tions of any agreement between the
25	parties, and, notwithstanding sub-

1	paragraph (A)(ii), may distribute such
2	returned product without providing
3	the transaction history. For trans-
4	actions subsequent to the return, the
5	transaction history of such product
6	shall begin with the wholesale dis-
7	tributor that accepted the returned
8	product, consistent with the require-
9	ments of this subsection.
10	"(II) Enhanced require-
11	MENTS.—Beginning 6 years after the
12	date of enactment of the Drug Supply
13	Chain Security Act (except as pro-
14	vided pursuant to subsection (a)(5)),
15	a wholesale distributor may accept re-
16	turned product from a dispenser or
17	repackager only if the wholesale dis-
18	tributor can associate returned prod-
19	uct with the transaction information
20	and transaction statement associated
21	with that product. For all trans-
22	actions after such date, the trans-
23	action history, as applicable, of such
24	product shall begin with the wholesale
25	distributor that accepted and verified

1	the returned product. For purposes of
2	this subparagraph, the transaction in-
3	formation and transaction history, as
4	applicable, need not include trans-
5	action dates if it is not reasonably
6	practicable to obtain such dates.
7	"(ii) Nonsaleable returns.—A
8	wholesale distributor may return a non-
9	saleable product to the manufacturer or re-
10	packager, to the wholesale distributor from
11	whom such product was purchased, or to a
12	person acting on behalf of such a person,
13	including a returns processor, without pro-
14	viding the information required under sub-
15	paragraph (A)(i).
16	"(C) Requests for information.—
17	Upon a request by the Secretary or other ap-
18	propriate Federal or State official, in the event
19	of a recall or for the purpose of investigating a
20	suspect product or an illegitimate product, a
21	wholesale distributor shall, not later than 1
22	business day, and not to exceed 48 hours, after
23	receiving the request or in other such reason-
24	able time as determined by the Secretary, based
25	on the circumstances of the request, provide the

1	applicable transaction information, transaction
2	history, and transaction statement for the prod-
3	$\operatorname{uct}$ .
4	"(D) Trading Partner agreements.—
5	Beginning 6 years after the date of enactment
6	of the Drug Supply Chain Security Act, a
7	wholesale distributor may disclose the trans-
8	action information, including lot level informa-
9	tion, transaction history, or transaction state-
10	ment of a product to the subsequent purchaser
11	of the product, pursuant to a written agreement
12	between such wholesale distributor and such
13	subsequent purchaser. Nothing in this subpara-
14	graph shall be construed to limit the applica-
15	bility of subparagraphs (A) through (C).
16	"(2) Product identifier.—Beginning 6
17	years after the date of enactment of the Drug Sup-
18	ply Chain Security Act, a wholesale distributor may
19	engage in transactions involving a product only if
20	such product is encoded with a product identifier
21	(except as provided pursuant to subsection (a)(5)).
22	"(3) Authorized trading partners.—Be-
23	ginning not later than January 1, 2015, the trading
24	partners of a wholesale distributor may be only au-
25	thorized trading partners.

1	"(4) Verification.—Beginning not later than
2	January 1, 2015, a wholesale distributor shall have
3	systems in place to enable the wholesale distributor
4	to comply with the following requirements:
5	"(A) Suspect product.—
6	"(i) In general.—Upon making a
7	determination that a product in the posses-
8	sion or control of a wholesale distributor is
9	a suspect product, or upon receiving a re-
10	quest for verification from the Secretary
11	that has made a determination that a
12	product within the possession or control of
13	a wholesale distributor is a suspect prod-
14	uct, a wholesale distributor shall—
15	"(I) quarantine such product
16	within the possession or control of the
17	wholesale distributor from product in-
18	tended for distribution until such
19	product is cleared or dispositioned;
20	and
21	"(II) promptly conduct an inves-
22	tigation in coordination with trading
23	partners, as applicable, to determine
24	whether the product is an illegitimate
25	product, which shall include validating

1	any applicable transaction history and
2	transaction information in the posses-
3	sion of the wholesale distributor and
4	otherwise investigating to determine
5	whether the product is an illegitimate
6	product, and, beginning 6 years after
7	the date of enactment of the Drug
8	Supply Chain Security Act (except as
9	provided pursuant to subsection
10	(a)(5)), verifying the product at the
11	package level, including the standard-
12	ized numerical identifier.
13	"(ii) CLEARED PRODUCT.—If the
14	wholesale distributor determines that a
15	suspect product is not an illegitimate prod-
16	uct, the wholesale distributor shall prompt-
17	ly notify the Secretary, if applicable, of
18	such determination and such product may
19	be further distributed.
20	"(iii) Records.—A wholesale dis-
21	tributor shall keep records of the investiga-
22	tion of a suspect product for not less than
23	6 years after the conclusion of the inves-
24	tigation.
25	"(B) Illegitimate product.—

1	"(i) In General.—Upon deter-
2	mining, in coordination with the manufac-
3	turer, that a product in the possession or
4	control of a wholesale distributor is an ille-
5	gitimate product, the wholesale distributor
6	shall, in a manner that is consistent with
7	the systems and processes of such whole-
8	sale distributor—
9	"(I) quarantine such product
10	within the possession or control of the
11	wholesale distributor from product in-
12	tended for distribution until such
13	product is dispositioned;
14	"(II) disposition the illegitimate
15	product within the possession or con-
16	trol of the wholesale distributor;
17	"(III) take reasonable and appro-
18	priate steps to assist a trading part-
19	ner to disposition an illegitimate prod-
20	uct not in the possession or control of
21	the wholesale distributor; and
22	"(IV) retain a sample of the
23	product for further physical examina-
24	tion or laboratory analysis of the
25	product by the manufacturer or Sec-

1	retary (or other appropriate Federal
2	or State official) upon request by the
3	manufacturer or Secretary (or other
4	appropriate Federal or State official),
5	as necessary and appropriate.
6	"(ii) Making a notification.—
7	Upon determining that a product in the
8	possession or control of the wholesale dis-
9	tributor is an illegitimate product, the
10	wholesale distributor shall notify the Sec-
11	retary and all immediate trading partners
12	that the wholesale distributor has reason
13	to believe may have received such illegit-
14	imate product of such determination not
15	later than 24 hours after making such de-
16	termination.
17	"(iii) Responding to a notifica-
18	TION.—Upon the receipt of a notification
19	from the Secretary or a trading partner
20	that a determination has been made that a
21	product is an illegitimate product, a whole-
22	sale distributor shall identify all illegit-
23	imate product subject to such notification
24	that is in the possession or control of the
25	wholesale distributor, including any prod-

1	uct that is subsequently received, and shall
2	perform the activities described in subpara-
3	graph (A).
4	"(iv) Terminating a notifica-
5	TION.—Upon making a determination, in
6	consultation with the Secretary, that a no-
7	tification is no longer necessary, a whole-
8	sale distributor shall promptly notify im-
9	mediate trading partners that the whole-
10	sale distributor notified pursuant to clause
11	(ii) that such notification has been termi-
12	nated.
13	"(v) Records.—A wholesale dis-
14	tributor shall keep records of the disposi-
15	tion of an illegitimate product for not less
16	than 6 years after the conclusion of the
17	disposition.
18	"(C) Electronic database.—A whole-
19	sale distributor may satisfy the requirements of
20	this paragraph by developing a secure electronic
21	database or utilizing a secure electronic data-
22	base developed or operated by another entity.
23	The owner of such database shall establish the
24	requirements and processes to respond to re-
25	quests and may provide for data access to other

1	members of the pharmaceutical distribution
2	supply chain, as appropriate. The development
3	and operation of such a database shall not re-
4	lieve a wholesale distributor of the requirement
5	under this paragraph to respond to a
6	verification request submitted by means other
7	than a secure electronic database.
8	"(D) VERIFICATION OF SALEABLE RE-
9	TURNED PRODUCT.—Beginning 6 years after
10	the date of enactment of the Drug Supply
11	Chain Security Act, upon receipt of a returned
12	product that the wholesale distributor intends
13	to further distribute, before further distributing
14	such product, the wholesale distributor shall
15	verify the product identifier, including the
16	standardized numerical identifier, for each
17	sealed homogeneous case of such product or, if
18	such product is not in a sealed homogeneous
19	case, verify the product identifier, including the
20	standardized numerical identifier, on each pack-
21	age.
22	"(d) Dispenser Requirements.—
23	"(1) Product tracing.—
24	"(A) In General.—Beginning July 1,
25	2015, a dispenser—

1	"(i) shall not accept ownership of a
2	product, unless the previous owner prior
3	to, or at the time of, the transaction, pro-
4	vides transaction history, transaction infor-
5	mation, and a transaction statement;
6	"(ii) prior to, or at the time of, each
7	transaction in which the dispenser trans-
8	fers ownership of a product (but not in-
9	cluding dispensing to a patient or returns)
10	shall provide the subsequent owner with
11	transaction history, transaction informa-
12	tion, and a transaction statement for the
13	product, except that the requirements of
14	this clause shall not apply to sales by a
15	dispenser to another dispenser to fulfill a
16	specific patient need; and
17	"(iii) shall capture transaction infor-
18	mation (including lot level information, if
19	provided), transaction history, and trans-
20	action statements, as necessary to inves-
21	tigate a suspect product, and maintain
22	such information, history, and statements
23	for not less than 6 years after the trans-
24	action.

1	"(B) AGREEMENTS WITH THIRD PAR-
2	TIES.—A dispenser may enter into a written
3	agreement with a third party, including an au-
4	thorized wholesale distributor, under which the
5	third party confidentially maintains the trans-
6	action information, transaction history, and
7	transaction statements required to be main-
8	tained under this subsection on behalf of the
9	dispenser. If a dispenser enters into such an
10	agreement, the dispenser shall maintain a copy
11	of the written agreement and shall not be re-
12	lieved of the obligations of the dispenser under
13	this subsection.
14	"(C) Returns.—
15	"(i) Saleable returns.—A dis-
16	penser may return product to the trading
17	partner from which the dispenser obtained
18	the product without providing the informa-
19	tion required under subparagraph (A).
20	"(ii) Nonsaleable returns.—A
21	dispenser may return a nonsaleable prod-
22	uct to the manufacturer or repackager, to
23	the wholesale distributor from whom such
24	product was purchased, to a returns proc-
25	essor, or to a person acting on behalf of

1	such a person without providing the infor-
2	mation required under subparagraph (A).
3	"(D) REQUESTS FOR INFORMATION.—
4	Upon a request by the Secretary or other ap-
5	propriate Federal or State official, in the event
6	of a recall or for the purpose of investigating a
7	suspect or an illegitimate product, a dispenser
8	shall, not later than 2 business days after re-
9	ceiving the request or in another such reason-
10	able time as determined by the Secretary, based
11	on the circumstances of the request, provide the
12	applicable transaction information, transaction
13	statement, and transaction history which the
14	dispenser received from the previous owner,
15	which shall not include the lot number of the
16	product, the initial transaction date, or the ini-
17	tial shipment date from the manufacturer un-
18	less such information was included in the trans-
19	action information, transaction statement, and
20	transaction history provided by the manufac-
21	turer or wholesale distributor to the dispenser.
22	The dispenser may respond to the request by
23	providing the applicable information in either
24	paper or electronic format. Until the date that
25	is 4 years after the date of enactment of the

1	Drug Supply Chain Security Act, the Secretary
2	or other appropriate Federal or State official
3	shall grant a dispenser additional time, as nec-
4	essary, only with respect to a request to provide
5	lot level information described in subparagraph
6	(F) of section 581(26) that was provided to the
7	dispenser in paper format, limit the request
8	time period to the 6 months preceding the re-
9	quest or other relevant date, and, in the event
10	of a recall, the Secretary, or other appropriate
11	Federal or State official may request informa-
12	tion only if such recall involves a serious ad-
13	verse health consequence or death to humans.
14	"(2) Product identifier.—Beginning not
15	later than 7 years after the date of enactment of the
16	Drug Supply Chain Security Act, a dispenser may
17	engage in transactions involving a product only if
18	such product is encoded with a product identifier
19	(except as provided pursuant to subsection (a)(5)).
20	"(3) Authorized trading partners.—Be-
21	ginning not later than January 1, 2015, the trading
22	partners of a dispenser may be only authorized trad-
23	ing partners.
24	"(4) Verification.—Beginning not later than
25	January 1, 2015, a dispenser shall have systems in

1	place to enable the dispenser to comply with the fol-
2	lowing requirements:
3	"(A) Suspect product.—
4	"(i) In General.—Upon making a
5	determination that a product in the posses-
6	sion or control of the dispenser is a suspect
7	product, or upon receiving a request for
8	verification from the Secretary that has
9	made a determination that a product with-
10	in the possession or control of a dispenser
11	is a suspect product, a dispenser shall—
12	"(I) quarantine such product
13	within the possession or control of the
14	dispenser from product intended for
15	distribution until such product is
16	cleared or dispositioned; and
17	"(II) promptly conduct an inves-
18	tigation in coordination with trading
19	partners, as applicable, to determine
20	whether the product is an illegitimate
21	product.
22	"(ii) Investigation.—An investiga-
23	tion conducted under clause (i)(II) shall in-
24	clude—

1	"(I) beginning 7 years after the
2	date of enactment of the Drug Supply
3	Chain Security Act, verifying whether
4	the lot number of a suspect product
5	corresponds with the lot number for
6	such product;
7	"(II) beginning 7 years after the
8	date of enactment of such Act,
9	verifying that the product identifier,
10	including the standardized numerical
11	identifier, of at least 3 packages or 10
12	percent of such suspect product,
13	whichever is greater, or all packages,
14	if there are fewer than 3, corresponds
15	with the product identifier for such
16	product;
17	"(III) validating any applicable
18	transaction history and transaction in-
19	formation in the possession of the dis-
20	penser; and
21	"(IV) otherwise investigating to
22	determine whether the product is an
23	illegitimate product.
24	"(iii) Cleared Product.—If the dis-
25	penser makes the determination that a sus-

1	pect product is not an illegitimate product,
2	the dispenser shall promptly notify the
3	Secretary, if applicable, of such determina-
4	tion and such product may be further dis-
5	tributed or dispensed.
6	"(iv) Records.—A dispenser shall
7	keep records of the investigation of a sus-
8	pect product for not less than 6 years after
9	the conclusion of the investigation.
10	"(B) Illegitimate product.—
11	"(i) In General.—Upon deter-
12	mining, in coordination with the manufac-
13	turer, that a product in the possession or
14	control of a dispenser is an illegitimate
15	product, the dispenser shall—
16	"(I) disposition the illegitimate
17	product within the possession or con-
18	trol of the dispenser;
19	"(II) take reasonable and appro-
20	priate steps to assist a trading part-
21	ner to disposition an illegitimate prod-
22	uct not in the possession or control of
23	the dispenser; and
24	"(III) retain a sample of the
25	product for further physical examina-

1	tion or laboratory analysis of the
2	product by the manufacturer or Sec-
3	retary (or other appropriate Federal
4	or State official) upon request by the
5	manufacturer or Secretary (or other
6	appropriate Federal or State official),
7	as necessary and appropriate.
8	"(ii) Making a notification.—
9	Upon determining that a product in the
10	possession or control of the dispenser is an
11	illegitimate product, the dispenser shall no-
12	tify the Secretary and all immediate trad-
13	ing partners that the dispenser has reason
14	to believe may have received such illegit-
15	imate product of such determination not
16	later than 24 hours after making such de-
17	termination.
18	"(iii) Responding to a notifica-
19	TION.—Upon the receipt of a notification
20	from the Secretary or a trading partner
21	that a determination has been made that a
22	product is an illegitimate product, a dis-
23	penser shall identify all illegitimate product
24	subject to such notification that is in the
25	possession or control of the dispenser, in-

1	cluding any product that is subsequently
2	received, and shall perform the activities
3	described in subparagraph (A).
4	"(iv) Terminating a notifica-
5	TION.—Upon making a determination, in
6	consultation with the Secretary, that a no-
7	tification is no longer necessary, a dis-
8	penser shall promptly notify immediate
9	trading partners that the dispenser notified
10	pursuant to clause (ii) that such notifica-
11	tion has been terminated.
12	"(v) Records.—A dispenser shall
13	keep records of the disposition of an illegit-
14	imate product for not less than 6 years
15	after the conclusion of the disposition.
16	"(C) Electronic database.—A dis-
17	penser may satisfy the requirements of this
18	paragraph by developing a secure electronic
19	database or utilizing a secure electronic data-
20	base developed or operated by another entity.
21	"(5) Exception.—Notwithstanding any other
22	provision of law, the requirements under paragraphs
23	(1) and (4) shall not apply to licensed health care
24	practitioners authorized to prescribe or administer
25	medication under State law or other licensed individ-

1	uals under the supervision or direction of such prac-
2	titioners who dispense or administer product in the
3	usual course of professional practice.
4	"(e) Repackager Requirements.—
5	"(1) Product tracing.—
6	"(A) In General.—Beginning not later
7	than January 1, 2015, a repackager described
8	in section 581(16)(A) shall—
9	"(i) not accept ownership of a product
10	unless the previous owner, prior to, or at
11	the time of, the transaction, provides
12	transaction history, transaction informa-
13	tion, and a transaction statement for the
14	product;
15	"(ii) prior to, or at the time of, each
16	transaction in which the repackager trans-
17	fers ownership of a product, provide the
18	subsequent owner with transaction history,
19	transaction information, and a transaction
20	statement for the product; and
21	"(iii) capture the transaction informa-
22	tion (including lot level information),
23	transaction history, and transaction state-
24	ment for each transaction described in
25	clauses (i) and (ii) and maintain such in-

1	formation, history, and statement for not
2	less than 6 years after the transaction.
3	"(B) Returns.—
4	"(i) Nonsaleable product.—A re-
5	packager described in section 581(16)(A)
6	may return a nonsaleable product to the
7	manufacturer or repackager, or to the
8	wholesale distributor from whom such
9	product was purchased, or to a person act-
10	ing on behalf of such a person, including
11	a returns processor, without providing the
12	information required under subparagraph
13	(A)(ii).
14	"(ii) Saleable or nonsaleable
15	PRODUCT.—A repackager described in sec-
16	tion 581(16)(B) may return a saleable or
17	nonsaleable product to the manufacturer,
18	repackager, or to the wholesale distributor
19	from whom such product was received
20	without providing the information required
21	under subparagraph (A)(ii) on behalf of
22	the hospital or other health care entity
23	that took ownership of such product pursu-
24	ant to the terms and conditions of any

1	agreement between such repackager and
2	the entity that owns the product.
3	"(C) Requests for information.—
4	Upon a request by the Secretary or other ap-
5	propriate Federal or State official, in the event
6	of a recall or for the purpose of investigating a
7	suspect product or an illegitimate product, a re-
8	packager described in section 581(16)(A) shall,
9	not later than 1 business day, and not to exceed
10	48 hours, after receiving the request or in other
11	such reasonable time as determined by the Sec-
12	retary, provide the applicable transaction infor-
13	mation, transaction history, and transaction
14	statement for the product.
15	"(2) Product identifier.—
16	"(A) IN GENERAL.—Beginning not later
17	than 5 years after the date of enactment of the
18	Drug Supply Chain Security Act, a repackager
19	described in section 581(16)(A)—
20	"(i) shall affix or imprint a product
21	identifier to each package and homogenous
22	case of product intended to be introduced
23	in a transaction in commerce;
24	"(ii) shall maintain the product iden-
25	tifier information for such product for not

1	less than 6 years after the date of the
2	transaction;
3	"(iii) may engage in transactions in-
4	volving a product only if such product is
5	encoded with a product identifier (except
6	as provided pursuant to subsection (a)(5));
7	and
8	"(iv) shall maintain records for not
9	less than 6 years to allow the repackager
10	to associate the product identifier the re-
11	packager affixes or imprints with the prod-
12	uct identifier assigned by the original man-
13	ufacturer of the product.
14	"(B) Exception.—A package that is re-
15	quired to have a standardized numerical identi-
16	fier is not required to have a unique device
17	identifier.
18	"(3) Authorized trading partners.—Be-
19	ginning January 1, 2015, the trading partners of a
20	repackager described in section 581(16) may be only
21	authorized trading partners.
22	"(4) Verification.—Beginning not later than
23	January 1, 2015, a repackager described in section
24	581(16)(A) shall have systems in place to enable the

1	repackager to comply with the following require-
2	ments:
3	"(A) Suspect product.—
4	"(i) In general.—Upon making a
5	determination that a product in the posses-
6	sion or control of the repackager is a sus-
7	pect product, or upon receiving a request
8	for verification from the Secretary that has
9	made a determination that a product with-
10	in the possession or control of a repack-
11	ager is a suspect product, a repackager
12	shall—
13	"(I) quarantine such product
14	within the possession or control of the
15	repackager from product intended for
16	distribution until such product is
17	cleared or dispositioned; and
18	"(II) promptly conduct an inves-
19	tigation in coordination with trading
20	partners, as applicable, to determine
21	whether the product is an illegitimate
22	product, which shall include validating
23	any applicable transaction history and
24	transaction information in the posses-
25	sion of the repackager and otherwise

1	investigating to determine whether the
2	product is an illegitimate product,
3	and, beginning 5 years after the date
4	of enactment of the Drug Supply
5	Chain Security Act (except as pro-
6	vided pursuant to subsection (a)(5)),
7	verifying the product at the package
8	level, including the standardized nu-
9	merical identifier.
10	"(ii) CLEARED PRODUCT.—If the re-
11	packager makes the determination that a
12	suspect product is not an illegitimate prod-
13	uct, the repackager shall promptly notify
14	the Secretary, if applicable, of such deter-
15	mination and such product may be further
16	distributed.
17	"(iii) Records.—A repackager shall
18	keep records of the investigation of a sus-
19	pect product for not less than 6 years after
20	the conclusion of the investigation.
21	"(B) Illegitimate product.—
22	"(i) In General.—Upon deter-
23	mining, in coordination with the manufac-
24	turer, that a product in the possession or
25	control of a repackager is an illegitimate

1	
1	product, the repackager shall, in a manner
2	that is consistent with the systems and
3	processes of such repackager—
4	"(I) quarantine such product
5	within the possession or control of the
6	repackager from product intended for
7	distribution until such product is
8	dispositioned;
9	"(II) disposition the illegitimate
10	product within the possession or con-
11	trol of the repackager;
12	"(III) take reasonable and appro-
13	priate steps to assist a trading part-
14	ner to disposition an illegitimate prod-
15	uct not in the possession or control of
16	the repackager; and
17	"(IV) retain a sample of the
18	product for further physical examina-
19	tion or laboratory analysis of the
20	product by the manufacturer or Sec-
21	retary (or other appropriate Federal
22	or State official) upon request by the
23	manufacturer or Secretary (or other
24	appropriate Federal or State official),
25	as necessary and appropriate.

1	"(ii) Making a notification.—
2	Upon determining that a product in the
3	possession or control of the repackager is
4	an illegitimate product, the repackager
5	shall notify the Secretary and all imme-
6	diate trading partners that the repackager
7	has reason to believe may have received the
8	illegitimate product of such determination
9	not later than 24 hours after making such
10	determination.
11	"(iii) Responding to a notifica-
12	TION.—Upon the receipt of a notification
13	from the Secretary or a trading partner, a
14	repackager shall identify all illegitimate
15	product subject to such notification that is
16	in the possession or control of the repack-
17	ager, including any product that is subse-
18	quently received, and shall perform the ac-
19	tivities described in subparagraph (A).
20	"(iv) TERMINATING A NOTIFICA-
21	TION.—Upon making a determination, in
22	consultation with the Secretary, that a no-
23	tification is no longer necessary, a repack-
24	ager shall promptly notify immediate trad-
25	ing partners that the repackager notified

1	pursuant to clause (ii) that such notifica-
2	tion has been terminated.
3	"(v) Records.—A repackager shall
4	keep records of the disposition of an illegit-
5	imate product for not less than 6 years
6	after the conclusion of the disposition.
7	"(C) Requests for Verification.—Be-
8	ginning 5 years after the date of enactment of
9	the Drug Supply Chain Security Act, upon re-
10	ceiving a request for verification from an au-
11	thorized manufacturer, wholesale distributor, or
12	dispenser that is in possession or control of a
13	product they believe to be repackaged by such
14	repackager, a repackager shall, not later than
15	24 hours after receiving the verification request
16	or in other such reasonable time as determined
17	by the Secretary, based on the circumstances of
18	the request, notify the person making the re-
19	quest whether the product identifier, including
20	the standardized numerical identifier, that is
21	the subject of the request corresponds to the
22	product identifier affixed or imprinted by the
23	repackager. If a repackager responding to a
24	verification request identifies a product identi-
25	fier that does not correspond to that affixed or

1	imprinted by the repackager, the repackager
2	shall treat such product as suspect product and
3	conduct an investigation as described in sub-
4	paragraph (A). If the repackager has reason to
5	believe the product is an illegitimate product
6	the repackager shall advise the person making
7	the request of such belief at the time such re-
8	packager responds to the verification request.
9	"(D) ELECTRONIC DATABASE.—A repack-
10	ager may satisfy the requirements of paragraph
11	(4) by developing a secure electronic database
12	or utilizing a secure electronic database devel-
13	oped or operated by another entity. The owner
14	of such database shall establish the require-
15	ments and processes to respond to requests and
16	may provide for data access to other members
17	of the pharmaceutical distribution supply chain
18	as appropriate. The development and operation
19	of such a database shall not relieve a repack-
20	ager of the requirement under subparagraph
21	(C) to respond to a verification request sub-
22	mitted by means other than a secure electronic
23	database.
24	"(E) VERIFICATION OF SALEABLE RE-
25	TURNED PRODUCT.—Beginning 5 years after

the date of enactment of the Drug Supply Chain Security Act, upon receipt of a returned product that the repackager intends to further distribute, before further distributing such product, the repackager shall verify the product identifier for each sealed homogeneous case of such product or, if such product is not in a sealed homogeneous case, verify the product identifier on each package.

## "(f) Drop Shipments.—

"(1) IN GENERAL.—A wholesale distributor that does not physically handle or store product shall be exempt from the provisions of this section, except the notification requirements under clauses (ii), (iii), and (iv) of subsection (c)(4)(B), provided that the manufacturer, repackager, or other wholesale distributor that distributes the product to the dispenser by means of a drop shipment for such wholesale distributor includes on the transaction information and transaction history to the dispenser the contact information of such wholesale distributor and provides the transaction information, transaction history, and transaction statement directly to the dispenser.

1	"(2) Clarification.—For purposes of this
2	subsection, providing administrative services, includ-
3	ing processing of orders and payments, shall not by
4	itself, be construed as being involved in the han-
5	dling, distribution, or storage of a product.".
6	SEC. 203. ENHANCED DRUG DISTRIBUTION SECURITY.
7	Section 582, as added by section 202, is amended by
8	adding at the end the following:
9	"(g) Enhanced Drug Distribution Security.—
10	"(1) In General.—On the date that is 10
11	years after the date of enactment of the Drug Sup-
12	ply Chain Security Act, the following interoperable,
13	electronic tracing of product at the package level re-
14	quirements shall go into effect:
15	"(A) The transaction information and the
16	transaction statements as required under this
17	section shall be exchanged in a secure, inter-
18	operable, electronic manner in accordance with
19	the standards established under the guidance
20	issued pursuant to paragraphs (3) and (4) of
21	subsection (h), including any revision of such
22	guidance issued in accordance with paragraph
23	(5) of such subsection.
24	"(B) The transaction information required
25	under this section shall include the product

1	identifier at the package level for each package
2	included in the transaction.
3	"(C) Systems and processes for verification
4	of product at the package level, including the
5	standardized numerical identifier, shall be re-
6	quired in accordance with the standards estab-
7	lished under the guidance issued pursuant to
8	subsection (a)(2) and the guidances issued pur-
9	suant to paragraphs (2), (3), and (4) of sub-
10	section (h), including any revision of such guid-
11	ances issued in accordance with paragraph (5)
12	of such subsection, which may include the use
13	of aggregation and inference as necessary.
14	"(D) The systems and processes necessary
15	to promptly respond with the transaction infor-
16	mation and transaction statement for a product
17	upon a request by the Secretary (or other ap-
18	propriate Federal or State official) in the event
19	of a recall or for the purposes of investigating
20	a suspect product or an illegitimate product
21	shall be required.
22	"(E) The systems and processes necessary
23	to promptly facilitate gathering the information
24	necessary to produce the transaction informa-

1	tion for each transaction going back to the
2	manufacturer, as applicable, shall be required—
3	"(i) in the event of a request by the
4	Secretary (or other appropriate Federal or
5	State official), on account of a recall or for
6	the purposes of investigating a suspect
7	product or an illegitimate product; or
8	"(ii) in the event of a request by an
9	authorized trading partner, in a secure
10	manner that ensures the protection of con-
11	fidential commercial information and trade
12	secrets, for purposes of investigating a sus-
13	pect product or assisting the Secretary (or
14	other appropriate Federal or State official)
15	with a request described in clause (i).
16	"(F) Each person accepting a saleable re-
17	turn shall have systems and processes in place
18	to allow acceptance of such product and may
19	accept saleable returns only if such person can
20	associate the saleable return product with the
21	transaction information and transaction state-
22	ment associated with that product.
23	"(2) Compliance.—
24	"(A) Information maintenance agree-
25	MENT.—A dispenser may enter into a written

1	agreement with a third party, including an au-
2	thorized wholesale distributor, under which the
3	third party shall confidentially maintain any in-
4	formation and statements required to be main-
5	tained under this section. If a dispenser enters
6	into such an agreement, the dispenser shall
7	maintain a copy of the written agreement and
8	shall not be relieved of the obligations of the
9	dispenser under this subsection.
10	"(B) ALTERNATIVE METHODS.—The Sec-
11	retary, taking into consideration the assessment
12	conducted under paragraph (3), shall provide
13	for alternative methods of compliance with any
14	of the requirements set forth in paragraph (1),
15	including—
16	"(i) establishing timelines for compli-
17	ance by small businesses (including small
18	business dispensers with 25 or fewer full-
19	time employees) with such requirements, in
20	order to ensure that such requirements do
21	not impose undue economic hardship for
22	small businesses, including small business
23	dispensers for whom the criteria set forth
24	in the assessment under paragraph (3) is
25	not met, if the Secretary determines that

1	such requirements under paragraph (1)
2	would result in undue economic hardship;
3	and
4	"(ii) establishing a process by which a
5	dispenser may request a waiver from any
6	of the requirements set forth in paragraph
7	(1) if the Secretary determines that such
8	requirements would result in an undue eco-
9	nomic hardship, which shall include a proc-
10	ess for the biennial review and renewal of
11	any such waiver.
12	"(3) Assessment.—
13	"(A) IN GENERAL.—Not later than the
14	date that is 18 months after the Secretary
15	issues the final guidance required under sub-
16	section (h), the Secretary shall enter into a con-
17	tract with a private, independent consulting
18	firm with expertise to conduct a technology and
19	software assessment that looks at the feasibility
20	of dispensers with 25 or fewer full-time employ-
21	ees conducting interoperable, electronic tracing
22	of products at the package level. Such assess-
23	ment shall be completed not later than $8\frac{1}{2}$
24	years after the date of enactment of the Drug
25	Supply Chain Security Act.

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1	"(B) Condition.—As a condition of the
2	award of the contract under subparagraph (A),
3	the private, independent consulting firm shall
4	agree to consult with dispensers with 25 or
5	fewer full-time employees when conducting the
6	assessment under such subparagraph.
7	"(C) CONTENT.—The assessment under
8	subparagraph (A) shall assess whether—
9	"(i) the necessary software and hard-
10	ware is readily accessible to such dis-
11	pensers;
12	"(ii) the necessary software and hard-
13	ware is prohibitively expensive to obtain,
14	install, and maintain for such dispensers;
15	and
16	"(iii) the necessary hardware and
17	software can be integrated into business
18	practices, such as interoperability with
19	wholesale distributors, for such dispensers.
20	"(D) Publication.—The Secretary
21	shall—
22	"(i) publish the statement of work for
23	the assessment under subparagraph (A)
24	for public comment prior to beginning the
25	assessment;

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1	"(ii) publish the final assessment for
2	public comment not later than 30 calendar
3	days after receiving such assessment; and
4	"(iii) hold a public meeting not later
5	than 180 calendar days after receiving the
6	final assessment at which public stake-
7	holders may present their views on the as-
8	sessment.
9	"(4) Procedure.—Notwithstanding section
10	553 of title 5, United States Code, the Secretary, in
11	promulgating any regulation pursuant to this sec-
12	tion, shall—
13	"(A) provide appropriate flexibility by—
14	"(i) not requiring the adoption of spe-
15	cific business systems for the maintenance
16	and transmission of data;
17	"(ii) prescribing alternative methods
18	of compliance for any of the requirements
19	set forth in paragraph (1) or set forth in
20	regulations implementing such require-
21	ments, including—
22	"(I) timelines for small busi-
23	nesses to comply with the require-
24	ments set forth in the regulations in
25	order to ensure that such require-

1	ments do not impose undue economic
2	hardship for small businesses (includ-
3	ing small business dispensers for
4	whom the criteria set forth in the as-
5	sessment under paragraph (3) is not
6	met), if the Secretary determines that
7	such requirements would result in
8	undue economic hardship; and
9	"(II) the establishment of a proc-
10	ess by which a dispenser may request
11	a waiver from any of the requirements
12	set forth in such regulations if the
13	Secretary determines that such re-
14	quirements would result in an undue
15	economic hardship; and
16	"(iii) taking into consideration—
17	"(I) the results of pilot projects,
18	including pilot projects pursuant to
19	this section and private sector pilot
20	projects, including those involving the
21	use of aggregation and inference;
22	"(II) the public meetings held
23	and related guidance documents
24	issued under this section;

1	"(III) the public health benefits
2	of any additional regulations in com-
3	parison to the cost of compliance with
4	such requirements, including on enti-
5	ties of varying sizes and capabilities;
6	"(IV) the diversity of the phar-
7	maceutical distribution supply chain
8	by providing appropriate flexibility for
9	each sector, including both large and
10	small businesses; and
11	"(V) the assessment pursuant to
12	paragraph (3) with respect to small
13	business dispensers, including related
14	public comment and the public meet-
15	ing, and requirements under this sec-
16	tion;
17	"(B) issue a notice of proposed rulemaking
18	that includes a copy of the proposed regulation;
19	"(C) provide a period of not less than 60
20	days for comments on the proposed regulation;
21	and
22	"(D) publish in the Federal Register the
23	final regulation not less than 2 years prior to
24	the effective date of the regulation.
25	"(h) Guidance Documents.—

1	"(1) In general.—For the purposes of facili-
2	tating the successful and efficient adoption of se-
3	cure, interoperable product tracing at the package
4	level in order to enhance drug distribution security
5	and further protect the public health, the Secretary
6	shall issue the guidance documents as provided for
7	in this subsection.
8	"(2) Suspect and illegitimate product.—
9	"(A) In General.—Not later than 180
10	days after the date of enactment of the Drug
11	Supply Chain Security Act, the Secretary shall
12	issue a guidance document to aid trading part-
13	ners in the identification of a suspect product
14	and notification termination. Such guidance
15	document shall—
16	"(i) identify specific scenarios that
17	could significantly increase the risk of a
18	suspect product entering the pharma-
19	ceutical distribution supply chain;
20	"(ii) provide recommendation on how
21	trading partners may identify such product
22	and make a determination on whether the
23	product is a suspect product as soon as
24	practicable; and

1	"(iii) set forth the process by which
2	manufacturers, repackagers, wholesale dis-
3	tributors, and dispensers shall terminate
4	notifications in consultation with the Sec-
5	retary regarding illegitimate product pur-
6	suant to subsections $(b)(4)(B)$ , $(c)(4)(B)$ ,
7	(d)(4)(B), and $(e)(4)(B)$ .
8	"(B) REVISED GUIDANCE.—If the Sec-
9	retary revises the guidance issued under sub-
10	paragraph (A), the Secretary shall follow the
11	procedure set forth in paragraph (5).
12	"(3) Unit level tracing.—
13	"(A) IN GENERAL.—In order to enhance
14	drug distribution security at the package level,
15	not later than 18 months after conducting a
16	public meeting on the system attributes nec-
17	essary to enable secure tracing of product at
18	the package level, including allowing for the use
19	of verification, inference, and aggregation, as
20	necessary, the Secretary shall issue a final guid-
21	ance document that outlines and makes rec-
22	ommendations with respect to the system at-
23	tributes necessary to enable secure tracing at
24	the package level as required under the require-

1	ments established under subsection (g). Such
2	guidance document shall—
3	"(i) define the circumstances under
4	which the sectors within the pharma-
5	ceutical distribution supply chain may, in
6	the most efficient manner practicable, infer
7	the contents of a case, pallet, tote, or other
8	aggregate of individual packages or con-
9	tainers of product, from a product identi-
10	fier associated with the case, pallet, tote,
11	or other aggregate, without opening each
12	case, pallet, tote, or other aggregate or
13	otherwise individually scanning each pack-
14	age;
15	"(ii) identify methods and processes
16	to enhance secure tracing of product at the
17	package level, such as secure processes to
18	facilitate the use of inference, enhanced
19	verification activities, the use of aggrega-
20	tion and inference, processes that utilize
21	the product identifiers to enhance tracing
22	of product at the package level, including
23	the standardized numerical identifier, or
24	package security features; and

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1	"(iii) ensure the protection of con-
2	fidential commercial information and trade
3	secrets.
4	"(B) Procedure.—In issuing the guid-
5	ance under subparagraph (A), and in revising
6	such guidance, if applicable, the Secretary shall
7	follow the procedure set forth in paragraph (5).
8	"(4) Standards for interoperable data
9	EXCHANGE.—
10	"(A) In General.—In order to enhance
11	secure tracing of a product at the package level,
12	the Secretary, not later than 18 months after
13	conducting a public meeting on the interoper-
14	able standards necessary to enhance the secu-
15	rity of the pharmaceutical distribution supply
16	chain, shall update the guidance issued pursu-
17	ant to subsection (a)(2), as necessary and ap-
18	propriate, and finalize such guidance document
19	so that the guidance document—
20	"(i) identifies and makes rec-
21	ommendations with respect to the stand-
22	ards necessary for adoption in order to
23	support the secure, interoperable electronic
24	data exchange among the pharmaceutical
25	distribution supply chain that comply with

1	a form and format developed by a widely
2	recognized international standards develop-
3	ment organization;
4	"(ii) takes into consideration stand-
5	ards established pursuant to subsection
6	(a)(2) and section 505D;
7	"(iii) facilitates the creation of a uni-
8	form process or methodology for product
9	tracing; and
10	"(iv) ensures the protection of con-
11	fidential commercial information and trade
12	secrets.
13	"(B) Procedure.—In issuing the guid-
14	ance under subparagraph (A), and in revising
15	such guidance, if applicable, the Secretary shall
16	follow the procedure set forth in paragraph (5).
17	"(5) Procedure.—In issuing or revising any
18	guidance issued pursuant to this subsection or sub-
19	section (g), except the initial guidance issued under
20	paragraph (2)(A), the Secretary shall—
21	"(A) publish a notice in the Federal Reg-
22	ister for a period not less than 30 days an-
23	nouncing that the draft or revised draft guid-
24	ance is available;

1	"(B) post the draft guidance document on
2	the Internet Web site of the Food and Drug
3	Administration and make such draft guidance
4	document available in hard copy;
5	"(C) provide an opportunity for comment
6	and review and take into consideration any
7	comments received;
8	"(D) revise the draft guidance, as appro-
9	priate;
10	"(E) publish a notice in the Federal Reg-
11	ister for a period not less than 30 days an-
12	nouncing that the final guidance or final revised
13	guidance is available;
14	"(F) post the final guidance document on
15	the Internet Web site of the Food and Drug
16	Administration and make such final guidance
17	document available in hard copy; and
18	"(G) provide for an effective date of not
19	earlier than 1 year after such guidance becomes
20	final.
21	"(i) Public Meetings.—
22	"(1) IN GENERAL.—The Secretary shall hold
23	not less than 5 public meetings to enhance the safe-
24	ty and security of the pharmaceutical distribution
25	supply chain and provide for comment. The Sec-

1	retary may hold the first such public meeting not
2	earlier than 1 year after the date of enactment of
3	the Drug Supply Chain Security Act. In carrying
4	out the public meetings described in this paragraph,
5	the Secretary shall—
6	"(A) prioritize topics necessary to inform
7	the issuance of the guidance described in para-
8	graphs (3) and (4) of subsection (h); and
9	"(B) take all measures reasonable and
10	practicable to ensure the protection of confiden-
11	tial commercial information and trade secrets.
12	"(2) Content.—Each of the following topics
13	shall be addressed in at least one of the public meet-
14	ings described in paragraph (1):
15	"(A) An assessment of the steps taken
16	under subsections (b) through (e) to build ca-
17	pacity for a unit-level system, including the im-
18	pact of the requirements of such subsections
19	on—
20	"(i) the ability of the health care sys-
21	tem collectively to maintain patient access
22	to medicines;
23	"(ii) the scalability of such require-
24	ments, including as it relates to product
25	lines; and

1	"(iii) the capability of different sec-
2	tors and subsectors, including both large
3	and small businesses, to affix and utilize
4	the product identifier.
5	"(B) The system attributes necessary to
6	support the requirements set forth under sub-
7	section (g), including the standards necessary
8	for adoption in order to support the secure,
9	interoperable electronic data exchange among
10	sectors within the pharmaceutical distribution
11	supply chain.
12	"(C) Best practices in each of the different
13	sectors within the pharmaceutical distribution
14	supply chain to implement the requirements of
15	this section.
16	"(D) The costs and benefits of the imple-
17	mentation of this section, including the impact
18	on each pharmaceutical distribution supply
19	chain sector and on public health.
20	"(E) Whether electronic tracing require-
21	ments, including tracing of product at the pack-
22	age level, are feasible, cost effective, and needed
23	to protect the public health.
24	"(F) The systems and processes needed to
25	utilize the product identifiers to enhance tracing

1	of product at the package level, including allow-
2	ing for verification, aggregation, and inference,
3	as necessary.
4	"(G) The technical capabilities and legal
5	authorities, if any, needed to establish an inter-
6	operable, electronic system that provides for
7	tracing of product at the package level.
8	"(H) The impact that such additional re-
9	quirements would have on patient safety, the
10	drug supply, cost and regulatory burden, and
11	timely patient access to prescription drugs.
12	"(I) Other topics, as determined appro-
13	priate by the Secretary.
14	"(j) Pilot Projects.—
15	"(1) IN GENERAL.—The Secretary shall estab-
16	lish 1 or more pilot projects, in coordination with
17	authorized manufacturers, repackagers, wholesale
18	distributors, and dispensers, to explore and evaluate
19	methods to enhance the safety and security of the
20	pharmaceutical distribution supply chain. Such
21	projects shall build upon efforts, in existence as of
22	the date of enactment of the Drug Supply Chain Se-
23	curity Act, to enhance the safety and security of the
24	pharmaceutical distribution supply chain, take into
25	consideration any pilot projects conducted prior to

1	such date of enactment, including any pilot projects
2	that use aggregation and inference, and inform the
3	draft and final guidance under paragraphs (3) and
4	(4) of subsection (h).
5	"(2) Content.—
6	"(A) IN GENERAL.—The Secretary shall
7	ensure that the pilot projects under paragraph
8	(1) reflect the diversity of the pharmaceutical
9	distribution supply chain and that the pilot
10	projects, when taken as a whole, include partici-
11	pants representative of every sector, including
12	both large and small businesses.
13	"(B) Project design.—The pilot
14	projects under paragraph (1) shall be designed
15	to—
16	"(i) utilize the product identifier for
17	tracing of a product, which may include
18	verification of the product identifier of a
19	product, including the use of aggregation
20	and inference;
21	"(ii) improve the technical capabilities
22	of each sector and subsector to comply
23	with systems and processes needed to uti-
24	lize the product identifiers to enhance trac-
25	ing of a product;

1	"(iii) identify system attributes that
2	are necessary to implement the require-
3	ments established under this section; and
4	"(iv) complete other activities as de-
5	termined by the Secretary.
6	"(k) Sunset.—The following requirements shall
7	have no force or effect beginning on the date that is 10
8	years after the date of enactment of the Drug Supply
9	Chain Security Act:
10	"(1) The provision and receipt of transaction
11	history under this section.
12	"(2) The requirements set forth for returns
13	$ under \qquad subsections \qquad (b)(4)(E), \qquad (e)(1)(B)(i),$
14	(d)(1)(C)(i), and $(e)(4)(E)$ .
15	"(3) The requirements set forth under subpara-
16	graphs $(A)(v)(II)$ and $(D)$ of subsection $(c)(1)$ , as
17	applied to lot level information only.
18	"(l) Rule of Construction.—The requirements
19	set forth in subsections (g)(4), (i), and (j) shall not be
20	construed as a condition, prohibition, or precedent for pre-
21	cluding or delaying the provisions becoming effective pur-
22	suant to subsection (g).
23	"(m) REQUESTS FOR INFORMATION.—On the date
24	that is 10 years after the date of enactment of the Drug
25	Supply Chain Security Act, the timeline for responses to

1	requests for information from the Secretary, or other ap-
2	propriate Federal or State official, as applicable, under
3	subsections (b)(1)(B), (c)(1)(C), and (e)(1)(C) shall be
4	not later than 24 hours after receiving the request from
5	the Secretary or other appropriate Federal or State offi-
6	cial, as applicable, or in such other reasonable time as de-
7	termined by the Secretary based on the circumstances of
8	the request.".
9	SEC. 204. NATIONAL STANDARDS FOR PRESCRIPTION DRUG
10	WHOLESALE DISTRIBUTORS.
11	(a) Amendments.—
12	(1) Requirement.—Section 503(e) (21 U.S.C.
13	353(e)) is amended by striking paragraphs (1), (2),
14	and (3) and inserting the following:
15	"(1) Requirement.—Subject to section 583:
16	"(A) In General.—No person may en-
17	gage in wholesale distribution of a drug subject
18	to subsection (b)(1) in any State unless such
19	person—
20	"(i)(I) is licensed by the State from
21	which the drug is distributed; or
22	"(II) if the State from which the drug
23	is distributed has not established a licen-
24	sure requirement, is licensed by the Sec-
25	retary; and

1	"(ii) if the drug is distributed inter-
2	state, is licensed by the State into which
3	the drug is distributed if the State into
4	which the drug is distributed requires the
5	licensure of a person that distributes drugs
6	into the State.
7	"(B) Standards.—Each Federal and
8	State license described in subparagraph (A)
9	shall meet the standards, terms, and conditions
10	established by the Secretary under section 583.
11	"(2) Reporting and database.—
12	"(A) Reporting.—Beginning January 1,
13	2015, any person who owns or operates an es-
14	tablishment that engages in wholesale distribu-
15	tion shall—
16	"(i) report to the Secretary, on an an-
17	nual basis pursuant to a schedule deter-
18	mined by the Secretary—
19	"(I) each State by which the per-
20	son is licensed and the appropriate
21	identification number of each such li-
22	cense; and
23	"(II) the name, address, and con-
24	tact information of each facility at
25	which, and all trade names under

1	which, the person conducts business;
2	and
3	"(ii) report to the Secretary within a
4	reasonable period of time and in a reason-
5	able manner, as determined by the Sec-
6	retary, any significant disciplinary actions,
7	such as the revocation or suspension of a
8	wholesale distributor license, taken by a
9	State or the Federal Government during
10	the reporting period against the wholesale
11	distributor.
12	"(B) Database.—Not later than January
13	1, 2015, the Secretary shall establish a data-
14	base of authorized wholesale distributors. Such
15	database shall—
16	"(i) identify each authorized wholesale
17	distributor by name, contact information,
18	and each State where such wholesale dis-
19	tributor is appropriately licensed to engage
20	in wholesale distribution;
21	"(ii) be available to the public on the
22	Internet Web site of the Food and Drug
23	Administration; and
24	"(iii) be regularly updated on a sched-
25	ule determined by the Secretary.

1	"(C) COORDINATION.—The Secretary shall
2	establish a format and procedure for appro-
3	priate State officials to access the information
4	provided pursuant to subparagraph (A) in a
5	prompt and secure manner.
6	"(D) Confidentiality.—Nothing in this
7	paragraph shall be construed as authorizing the
8	Secretary to disclose any information that is a
9	trade secret or confidential information subject
10	to section 552(b)(4) of title 5, United States
11	Code, or section 1905 of title 18, United States
12	Code.
13	"(3) Costs.—
14	"(A) AUTHORIZED FEES OF SECRETARY.—
15	If a State does not establish a licensing pro-
16	gram for persons engaged in the wholesale dis-
17	tribution of a drug subject to subsection (b),
18	the Secretary shall license a person engaged in
19	wholesale distribution located in such State and
20	may collect a reasonable fee in such amount
21	necessary to reimburse the Secretary for costs
22	associated with establishing and administering
23	the licensure program and conducting periodic
24	inspections under this section. The Secretary
25	shall adjust fee rates as needed on an annual

1	basis to generate only the amount of revenue
2	needed to perform this service. Fees authorized
3	under this paragraph shall be collected and
4	available for obligation only to the extent and in
5	the amount provided in advance in appropria-
6	tions Acts. Such fees are authorized to remain
7	available until expended. Such sums as may be
8	necessary may be transferred from the Food
9	and Drug Administration salaries and expenses
10	appropriation account without fiscal year limi-
11	tation to such appropriation account for sala-
12	ries and expenses with such fiscal year limita-
13	tion.
14	"(B) STATE LICENSING FEES.—Nothing in
15	this Act shall prohibit States from collecting
16	fees from wholesale distributors in connection
17	with State licensing of such distributors.".
18	(2) Wholesale distribution.—Section
19	503(e) (21 U.S.C. 353(e)), as amended by para-
20	graph (1), is further amended by adding at the end
21	the following:
22	"(4) For the purposes of this subsection and
23	subsection (d), the term 'wholesale distribution'
24	means the distribution of a drug subject to sub-
25	section (b) to a person other than a consumer or pa-

1	tient, or receipt of a drug subject to subsection (b)
2	by a person other than the consumer or patient, but
3	does not include—
4	"(A) intracompany distribution of any
5	drug between members of an affiliate or within
6	a manufacturer;
7	"(B) the distribution of a drug, or an offer
8	to distribute a drug among hospitals or other
9	health care entities which are under common
10	control;
11	"(C) the distribution of a drug or an offer
12	to distribute a drug for emergency medical rea-
13	sons, including a public health emergency dec-
14	laration pursuant to section 319 of the Public
15	Health Service Act, except that, for purposes of
16	this paragraph, a drug shortage not caused by
17	a public health emergency shall not constitute
18	an emergency medical reason;
19	"(D) the dispensing of a drug pursuant to
20	a prescription executed in accordance with sub-
21	section (b)(1);
22	"(E) the distribution of minimal quantities
23	of drug by a licensed retail pharmacy to a li-
24	censed practitioner for office use:

1	"(F) the distribution of a drug or an offer
2	to distribute a drug by a charitable organization
3	to a nonprofit affiliate of the organization to
4	the extent otherwise permitted by law;
5	"(G) the purchase or other acquisition by
6	a dispenser, hospital, or other health care entity
7	of a drug for use by such dispenser, hospital, or
8	other health care entity;
9	"(H) the distribution of a drug by the
10	manufacturer of such drug;
11	"(I) the receipt or transfer of a drug by an
12	authorized third-party logistics provider pro-
13	vided that such third-party logistics provider
14	does not take ownership of the drug;
15	"(J) a common carrier that transports a
16	drug, provided that the common carrier does
17	not take ownership of the drug;
18	"(K) the distribution of a drug, or an offer
19	to distribute a drug by an authorized repack-
20	ager that has taken ownership or possession of
21	the drug and repacks it in accordance with sec-
22	tion 582(e);
23	"(L) salable drug returns when conducted
24	by a dispenser;

1	"(M) the distribution of a collection of fin-
2	ished medical devices, which may include a
3	product or biological product, assembled in kit
4	form strictly for the convenience of the pur-
5	chaser or user (referred to in this subparagraph
6	as a 'medical convenience kit') if—
7	"(i) the medical convenience kit is as-
8	sembled in an establishment that is reg-
9	istered with the Food and Drug Adminis-
10	tration as a device manufacturer in accord-
11	ance with section $510(b)(2)$ ;
12	"(ii) the medical convenience kit does
13	not contain a controlled substance that ap-
14	pears in a schedule contained in the Com-
15	prehensive Drug Abuse Prevention and
16	Control Act of 1970;
17	"(iii) in the case of a medical conven-
18	ience kit that includes a product, the per-
19	son that manufacturers the kit—
20	"(I) purchased such product di-
21	rectly from the pharmaceutical manu-
22	facturer or from a wholesale dis-
23	tributor that purchased the product
24	directly from the pharmaceutical man-
25	ufacturer; and

1	"(II) does not alter the primary
2	container or label of the product as
3	purchased from the manufacturer or
4	wholesale distributor; and
5	"(iv) in the case of a medical conven-
6	ience kit that includes a product, the prod-
7	uct is—
8	"(I) an intravenous solution in-
9	tended for the replenishment of fluids
10	and electrolytes;
11	"(II) a product intended to main-
12	tain the equilibrium of water and min-
13	erals in the body;
14	"(III) a product intended for irri-
15	gation or reconstitution;
16	"(IV) an anesthetic;
17	"(V) an anticoagulant;
18	"(VI) a vasopressor; or
19	"(VII) a sympathomimetic;
20	"(N) the distribution of an intravenous
21	drug that, by its formulation, is intended for
22	the replenishment of fluids and electrolytes
23	(such as sodium, chloride, and potassium) or
24	calories (such as dextrose and amino acids);

1	"(O) the distribution of an intravenous
2	drug used to maintain the equilibrium of water
3	and minerals in the body, such as dialysis solu-
4	tions;
5	"(P) the distribution of a drug that is in-
6	tended for irrigation, or sterile water, whether
7	intended for such purposes or for injection;
8	"(Q) the distribution of medical gas, as de-
9	fined in section 575;
10	"(R) facilitating the distribution of a prod-
11	uct by providing solely administrative services,
12	including processing of orders and payments; or
13	"(S) the transfer of a product by a hos-
14	pital or other health care entity, or by a whole-
15	sale distributor or manufacturer operating at
16	the direction of the hospital or other health care
17	entity, to a repackager described in section
18	581(16)(B) and registered under section 510
19	for the purpose of repackaging the drug for use
20	by that hospital, or other health care entity and
21	other health care entities that are under com-
22	mon control, if ownership of the drug remains
23	with the hospital or other health care entity at
24	all times.".

1	(3) Third-party logistics providers.—Sec-
2	tion 503(e) (21 U.S.C. 353(e)), as amended by para-
3	graph (2), is further amended by adding at the end
4	the following:
5	"(5) Third-party logistics providers.—
6	Notwithstanding paragraphs (1) through (4), each
7	entity that meets the definition of a third-party lo-
8	gistics provider under section 581(22) shall obtain a
9	license as a third-party logistics provider as de-
10	scribed in section 584(a) and is not required to ob-
11	tain a license as a wholesale distributor if the entity
12	never assumes an ownership interest in the product
13	it handles.".
14	(4) Affiliate.—Section 503(e) (21 U.S.C.
15	353(e)), as amended by paragraph (3), is further
16	amended by adding at the end the following:
17	"(6) Affiliate.—For purposes of this sub-
18	section, the term 'affiliate' means a business entity
19	that has a relationship with a second business entity
20	if, directly or indirectly—
21	"(A) one business entity controls, or has
22	the power to control, the other business entity;
23	or
24	"(B) a third party controls, or has the
25	power to control, both of the business entities.".

1	(5) STANDARDS.—Subchapter H of chapter V,
2	as added by section 202, is amended by adding at
3	the end the following:
4	"SEC. 583. NATIONAL STANDARDS FOR PRESCRIPTION
5	DRUG WHOLESALE DISTRIBUTORS.
6	"(a) In General.—The Secretary shall, not later
7	than 2 years after the date of enactment of the Drug Sup-
8	ply Chain Security Act, establish by regulation standards
9	for the licensing of persons under section 503(e)(1) (as
10	amended by the Drug Supply Chain Security Act), includ-
11	ing the revocation, reissuance, and renewal of such license.
12	"(b) Content.—For the purpose of ensuring uni-
13	formity with respect to standards set forth in this section,
14	the standards established under subsection (a) shall apply
15	to all State and Federal licenses described under section
16	503(e)(1) (as amended by the Drug Supply Chain Secu-
17	rity Act) and shall include standards for the following:
18	"(1) The storage and handling of prescription
19	drugs, including facility requirements.
20	"(2) The establishment and maintenance of
21	records of the distributions of such drugs.
22	"(3) The furnishing of a bond or other equiva-
23	lent means of security, as follows:
24	"(A)(i) For the issuance or renewal of a
25	wholesale distributor license, an applicant that

1	is not a government owned and operated whole-
2	sale distributor shall submit a surety bond of
3	\$100,000 or other equivalent means of security
4	acceptable to the State.
5	"(ii) For purposes of clause (i), the State
6	or other applicable authority may accept a sur-
7	ety bond in the amount of \$25,000 if the an-
8	nual gross receipts of the previous tax year for
9	the wholesaler is \$10,000,000 or less.
10	"(B) If a wholesale distributor can provide
11	evidence that it possesses the required bond in
12	a State, the requirement for a bond in another
13	State shall be waived.
14	"(4) Mandatory background checks and
15	fingerprinting of facility managers or designated
16	representatives.
17	"(5) The establishment and implementation of
18	qualifications for key personnel.
19	"(6) The mandatory physical inspection of any
20	facility to be used in wholesale distribution within a
21	reasonable time frame from the initial application of
22	the facility and to be conducted by the licensing au-
23	thority or by the State, consistent with subsection
24	(e).

1	"(7) In accordance with subsection (d), the pro-
2	hibition of certain persons from receiving or main-
3	taining licensure for wholesale distribution.
4	"(c) Inspections.—To satisfy the inspection re-
5	quirement under subsection (b)(6), the Federal or State
6	licensing authority may conduct the inspection or may ac-
7	cept an inspection by the State in which the facility is lo-
8	cated, or by a third-party accreditation or inspection serv-
9	ice approved by the Secretary or the State licensing such
10	wholesale distributor.
11	"(d) Prohibited Persons.—The standards estab-
12	lished under subsection (a) shall include requirements to
13	prohibit a person from receiving or maintaining licensure
14	for wholesale distribution if the person—
15	"(1) has been convicted of any felony for con-
16	duct relating to wholesale distribution, any felony
17	violation of subsection (i) or (k) of section 301, or
18	any felony violation of section 1365 of title 18,
19	United States Code, relating to product tampering;
20	or
21	"(2) has engaged in a pattern of violating the
22	requirements of this section, or State requirements
23	for licensure, that presents a threat of serious ad-
24	verse health consequences or death to humans.

1	"(e) REQUIREMENTS.—The Secretary, in promul-
2	gating any regulation pursuant to this section, shall, not-
3	withstanding section 553 of title 5, United States Code—
4	"(1) issue a notice of proposed rulemaking that
5	includes a copy of the proposed regulation;
6	"(2) provide a period of not less than 60 days
7	for comments on the proposed regulation; and
8	"(3) provide that the final regulation take effect
9	on the date that is 2 years after the date such final
10	regulation is published.".
11	(b) Authorized Distributors of Record.—Sec-
12	tion 503(d) (21 U.S.C. 353(d)) is amended by adding at
13	the end the following:
14	"(4) In this subsection, the term 'authorized
15	distributors of record' means those distributors with
16	whom a manufacturer has established an ongoing re-
17	lationship to distribute such manufacturer's prod-
18	ucts.".
19	(c) Effective Date.—The amendments made by
20	subsections (a) and (b) shall take effect on January 1,
21	2015.

1	SEC. 205. NATIONAL STANDARDS FOR THIRD-PARTY LOGIS-
2	TICS PROVIDERS; UNIFORM NATIONAL POL-
3	ICY.
4	Subchapter H of chapter V, as amended by section
5	204, is further amended by adding at the end the fol-
6	lowing:
7	"SEC. 584. NATIONAL STANDARDS FOR THIRD-PARTY LO-
8	GISTICS PROVIDERS.
9	"(a) Requirements.—No third-party logistics pro-
10	vider in any State may conduct activities in any State un-
11	less each facility of such third-party logistics provider—
12	"(1)(A) is licensed by the State from which the
13	drug is distributed by the third-party logistics pro-
14	vider, in accordance with the regulations promul-
15	gated under subsection (d); or
16	"(B) if the State from which the drug distrib-
17	uted by the third-party logistics provider has not es-
18	tablished a licensure requirement, is licensed by the
19	Secretary, in accordance with the regulations pro-
20	mulgated under subsection (d); and
21	"(2) if the drug is distributed interstate, is li-
22	censed by the State into which the drug is distrib-
23	uted by the third-party logistics provider if such
24	State licenses third-party logistics providers that dis-
25	tribute drugs into the State and the third-party lo-

1	gistics provider is not licensed by the Secretary as
2	described in paragraph (1)(B).
3	"(b) Reporting.—Beginning 1 year after the date
4	of enactment of the Drug Supply Chain Security Act, a
5	facility of a third-party logistics provider shall report to
6	the Secretary, on an annual basis pursuant to a schedule
7	determined by the Secretary—
8	"(1) the State by which the facility is licensed
9	and the appropriate identification number of such li-
10	cense; and
11	"(2) the name and address of the facility and
12	all trade names under which such facility conducts
13	business.
14	"(c) Costs.—
15	"(1) Authorized fees of secretary.—If a
16	State does not establish a licensing program for a
17	third-party logistics provider, the Secretary shall li-
18	cense the third-party logistics provider located in
19	such State and may collect a reasonable fee in such
20	amount necessary to reimburse the Secretary for
21	costs associated with establishing and administering
22	the licensure program and conducting periodic in-
23	spections under this section. The Secretary shall ad-
24	just fee rates as needed on an annual basis to gen-
25	erate only the amount of revenue needed to perform

1	this service. Fees authorized under this paragraph
2	shall be collected and available for obligation only to
3	the extent and in the amount provided in advance in
4	appropriations Acts. Such fees are authorized to re-
5	main available until expended. Such sums as may be
6	necessary may be transferred from the Food and
7	Drug Administration salaries and expenses appro-
8	priation account without fiscal year limitation to
9	such appropriation account for salaries and expenses
10	with such fiscal year limitation.
11	"(2) State licensing fees.—
12	"(A) State established program.—
13	Nothing in this Act shall prohibit a State that
14	has established a program to license a third-
15	party logistics provider from collecting fees
16	from a third-party logistics provider for such a
17	license.
18	"(B) No state established pro-
19	GRAM.—A State that does not establish a pro-
20	gram to license a third-party logistics provider
21	in accordance with this section shall be prohib-
22	ited from collecting a State licensing fee from
23	a third-party logistics provider.
24	"(d) Regulations.—

1	"(1) In general.—Not later than 2 years
2	after the date of enactment of the Drug Supply
3	Chain Security Act, the Secretary shall issue regula-
4	tions regarding the standards for licensing under
5	subsection (a), including the revocation and
6	reissuance of such license, to third-party logistics
7	providers under this section.
8	"(2) Content.—Such regulations shall—
9	"(A) establish a process by which a third-
10	party accreditation program approved by the
11	Secretary shall, upon request by a third-party
12	logistics provider, issue a license to each third-
13	party logistics provider that meets the require-
14	ments set forth in this section;
15	"(B) establish a process by which the Sec-
16	retary shall issue a license to each third-party
17	logistics provider that meets the requirements
18	set forth in this section if the Secretary is not
19	able to approve a third-party accreditation pro-
20	gram because no such program meets the Sec-
21	retary's requirements necessary for approval of
22	such a third-party accreditation program;
23	"(C) require that the entity complies with
24	storage practices, as determined by the Sec-
25	retary for such facility, including—

1	"(i) maintaining access to warehouse
2	space of suitable size to facilitate safe op-
3	erations, including a suitable area to quar-
4	antine suspect product;
5	"(ii) maintaining adequate security;
6	and
7	"(iii) having written policies and pro-
8	cedures to—
9	"(I) address receipt, security,
10	storage, inventory, shipment, and dis-
11	tribution of a product;
12	"(II) identify, record, and report
13	confirmed losses or thefts in the
14	United States;
15	"(III) correct errors and inac-
16	curacies in inventories;
17	"(IV) provide support for manu-
18	facturer recalls;
19	"(V) prepare for, protect against,
20	and address any reasonably foresee-
21	able crisis that affects security or op-
22	eration at the facility, such as a
23	strike, fire, or flood;
24	"(VI) ensure that any expired
25	product is segregated from other

1	products and returned to the manu-
2	facturer or repackager or destroyed;
3	"(VII) maintain the capability to
4	trace the receipt and outbound dis-
5	tribution of a product, and supplies
6	and records of inventory; and
7	"(VIII) quarantine or destroy a
8	suspect product if directed to do so by
9	the respective manufacturer, wholesale
10	distributor, dispenser, or an author-
11	ized government agency;
12	"(D) provide for periodic inspection by the
13	licensing authority, as determined by the Sec-
14	retary, of such facility warehouse space to en-
15	sure compliance with this section;
16	"(E) prohibit a facility from having as a
17	manager or designated representative anyone
18	convicted of any felony violation of subsection
19	(i) or (k) of section 301 or any violation of sec-
20	tion 1365 of title 18, United States Code relat-
21	ing to product tampering;
22	"(F) provide for mandatory background
23	checks of a facility manager or a designated
24	representative of such manager;

1	"(G) require a third-party logistics pro-
2	vider to provide the applicable licensing author-
3	ity, upon a request by such authority, a list of
4	all product manufacturers, wholesale distribu-
5	tors, and dispensers for whom the third-party
6	logistics provider provides services at such facil-
7	ity; and
8	"(H) include procedures under which any
9	third-party logistics provider license—
10	"(i) expires on the date that is 3
11	years after issuance of the license; and
12	"(ii) may be renewed for additional 3-
13	year periods.
14	"(3) Procedure.—In promulgating the regula-
15	tions under this subsection, the Secretary shall, not-
16	withstanding section 553 of title 5, United States
17	Code—
18	"(A) issue a notice of proposed rulemaking
19	that includes a copy of the proposed regulation;
20	"(B) provide a period of not less than 60
21	days for comments on the proposed regulation;
22	and
23	"(C) provide that the final regulation takes
24	effect upon the expiration of 1 year after the
25	date that such final regulation is issued.

- 1 "(e) Validity.—A license issued under this section
- 2 shall remain valid as long as such third-party logistics pro-
- 3 vider remains licensed consistent with this section. If the
- 4 Secretary finds that the third-party accreditation program
- 5 demonstrates that all applicable requirements for licensure
- 6 under this section are met, the Secretary shall issue a li-
- 7 cense under this section to a third-party logistics provider
- 8 receiving accreditation, pursuant to subsection (d)(2)(A).

## 9 "SEC. 585. UNIFORM NATIONAL POLICY.

- 10 "(a) Product Tracing and Other Require-
- 11 MENTS.—Beginning on the date of enactment of the Drug
- 12 Supply Chain Security Act, no State or political subdivi-
- 13 sion of a State may establish or continue in effect any
- 14 requirements for tracing products through the distribution
- 15 system (including any requirements with respect to state-
- 16 ments of distribution history, transaction history, trans-
- 17 action information, or transaction statement of a product
- 18 as such product changes ownership in the supply chain,
- 19 or verification, investigation, disposition, notification, or
- 20 recordkeeping relating to such systems, including paper or
- 21 electronic pedigree systems or for tracking and tracing
- 22 drugs throughout the distribution system) which are in-
- 23 consistent with, more stringent than, or in addition to, any
- 24 requirements applicable under section 503(e) (as amended

1	by such Act) or this subchapter (or regulations issued
2	thereunder), or which are inconsistent with—
3	"(1) any waiver, exception, or exemption pursu-
4	ant to section 581 or 582; or
5	"(2) any restrictions specified in section 582.
6	"(b) Wholesale Distributor and Third-Party
7	LOGISTICS PROVIDER STANDARDS.—
8	"(1) In general.—Beginning on the date of
9	enactment of the Drug Supply Chain Security Act,
10	no State or political subdivision of a State may es-
11	tablish or continue any standards, requirements, or
12	regulations with respect to wholesale prescription
13	drug distributor or third-party logistics provider li-
14	censure that are inconsistent with, less stringent
15	than, directly related to, or covered by the standards
16	and requirements applicable under section 503(e)
17	(as amended by such Act), in the case of a wholesale
18	distributor, or section 584, in the case of a third-
19	party logistics provider.
20	"(2) State regulation of third-party lo-
21	GISTICS PROVIDERS.—No State shall regulate third-
22	party logistics providers as wholesale distributors.
23	"(3) Administration fees.—Notwithstanding
24	paragraph (1), a State may administer fee collec-
25	tions for effectuating the wholesale drug distributor

1	and third-party logistics provider licensure require-
2	ments under sections 503(e) (as amended by the
3	Drug Supply Chain Security Act), 583, and 584.
4	"(4) Enforcement, suspension, and rev-
5	OCATION.—Notwithstanding paragraph (1), a
6	State—
7	"(A) may take administrative action, in-
8	cluding fines, to enforce a requirement promul-
9	gated by the State in accordance with section
10	503(e) (as amended by the Drug Supply Chain
11	Security Act) or this subchapter;
12	"(B) may provide for the suspension or
13	revocation of licenses issued by the State for
14	violations of the laws of such State;
15	"(C) upon conviction of violations of Fed-
16	eral, State, or local drug laws or regulations,
17	may provide for fines, imprisonment, or civil
18	penalties; and
19	"(D) may regulate activities of licensed en-
20	tities in a manner that is consistent with prod-
21	uct tracing requirements under section 582.
22	"(c) Exception.—Nothing in this section shall be
23	construed to preempt State requirements related to the
24	distribution of prescription drugs if such requirements are
25	not related to product tracing as described in subsection

- 1 (a) or wholesale distributor and third-party logistics pro-
- 2 vider licensure as described in subsection (b) applicable
- 3 under section 503(e) (as amended by the Drug Supply
- 4 Chain Security Act) or this subchapter (or regulations
- 5 issued thereunder).".
- 6 SEC. 206. PENALTIES.
- 7 (a) Prohibited Act.—Section 301(t) (21 U.S.C.
- 8 331(t)), is amended—
- 9 (1) by striking "or" after "the requirements of
- section 503(d),"; and
- 11 (2) by inserting ", failure to comply with the
- requirements under section 582, the failure to com-
- ply with the requirements under section 584, as ap-
- plicable," after "in violation of section 503(e)".
- 15 (b) MISBRANDING.—Section 502 (21 U.S.C. 352), as
- 16 amended by section 103, is further amended by adding
- 17 at the end the following:
- 18 "(cc) If it is a drug and it fails to bear the product
- 19 identifier as required by section 582.".
- 20 SEC. 207. CONFORMING AMENDMENT.
- 21 (a) IN GENERAL.—Section 303(b)(1)(D) (21 U.S.C.
- 22 333(b)(1)(D)) is amended by striking "503(e)(2)(A)" and
- 23 inserting "503(e)(1)".
- 24 (b) Effective Date.—The amendment made by
- 25 subsection (a) shall take effect on January 1, 2015.

## 1 SEC. 208. SAVINGS CLAUSE.

- 2 Except as provided in the amendments made by para-
- 3 graphs (1), (2), and (3) of section 204(a) and by section
- 4 206(a), nothing in this title (including the amendments
- 5 made by this title) shall be construed as altering any au-
- 6 thority of the Secretary of Health and Human Services
- 7 with respect to a drug subject to section 503(b)(1) of the
- 8 Federal Food, Drug, and Cosmetic Act (21 U.S.C.
- 9 353(b)(1)) under any other provision of such Act or the
- 10 Public Health Service Act (42 U.S.C. 201 et seq.).