To amend the Federal Food, Drug, and Cosmetic Act with respect to compounding drugs.

IN THE SENATE OF THE UNITED STATES

	introduced the following bill; which was read twi-	сe
and referred	to the Committee on	

A BILL

To amend the Federal Food, Drug, and Cosmetic Act with respect to compounding drugs.

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; REFERENCES IN ACT.
4	(a) Short Title.—This Act may be cited as the
5	["Act"].
6	(b) References in Act.—Except as otherwise spec-
7	ified, amendments made by this Act to a section or other
8	provision of law are amendments to such section or other
9	provision of the Federal Food, Drug, and Cosmetic Act

10 (21 U.S.C. 301 et seq.).

1	SEC. 2. REGULATION OF HUMAN AND ANIMAL DRUG
2	COMPOUNDING.
3	(a) Clarification of New Drug Status.—For
4	purposes of the Federal Food, Drug, and Cosmetic Act
5	(21 U.S.C. 301 et seq.), the term "new drug" (as defined
6	in section 201(p) of such Act) shall include a compounded
7	drug.
8	(b) REGULATION OF HUMAN AND ANIMAL DRUG
9	Compounding.—Section 503A (21 U.S.C. 353a) is
10	amended to read as follows:
11	"SEC. 503A. HUMAN AND ANIMAL DRUG COMPOUNDING.
12	"(a) Definitions.—In this section:
13	"(1) Compounding.—The terms
14	'compounding' and 'compound'—
15	"(A) include compounding from bulk sub-
16	stances, admixing, and repackaging; and
17	"(B) do not include mixing, reconstituting,
18	or other such acts that are performed in ac-
19	cordance with directions contained in approved
20	labeling provided by the product's manufac-
21	turer.
22	"(2) Compounding manufacturer.—
23	"(A) IN GENERAL.—The term
24	'compounding manufacturer' means an entity—
25	"(i) that compounds any sterile drug
26	without receiving a prescription order for

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1	such drug prior to beginning compounding,
2	and introduces such compounded drug into
3	interstate commerce; or
4	"(ii) that repackages a drug using
5	sterile preservative-free single-dose vials or
6	by pooling sterile drugs.
7	"(B) EXCLUDED ACTIVITIES.—An entity
8	shall not be considered a compounding manu-
9	facturer if such entity—
10	"(i) repackages drugs in accordance
11	with section 506F; and
12	"(ii) does not otherwise meet the defi-
13	nition of compounding manufacturer under
14	subparagraph (A).
15	"(3) Traditional compounder.—
16	"(A) IN GENERAL.—The term 'traditional
17	compounder' means an entity—
18	"(i) wherein a drug is compounded
19	by—
20	"(I) a licensed pharmacist in a
21	State-licensed pharmacy; or
22	"(II) a licensed physician or li-
23	censed veterinarian, to the extent per-
24	mitted under State law;
25	"(ii) that—

4

1	"(I) compounds a drug upon re-
2	ceipt of a prescription order for an
3	identified individual patient; or
4	"(II) compounds a drug in lim-
5	ited quantities before receipt of a pre-
6	scription order for an identified indi-
7	vidual patient if such compounding is
8	based on a history of the licensed
9	pharmacist, licensed physician, or li-
10	censed veterinarian receiving prescrip-
11	tion orders for the compounding of
12	the drug, which orders have been gen-
13	erated solely within an established re-
14	lationship between the licensed phar-
15	macist, licensed physician, or licensed
16	veterinarian and—
17	"(aa) such individual patient
18	for whom the prescription order
19	will be provided; or
20	"(bb) the licensed physician
21	licensed veterinarian, or other li-
22	censed practitioner who will write
23	such prescription order; and
24	"(iii) that does not perform any of the
25	activities described in clause (i) or (ii) of

1	paragraph (2)(A), except as provided in
2	subparagraph (B).
3	"(B) Exception.—A pharmacy within a
4	health system (as defined in section 506F) that
5	compounds a drug and ships such drug for dis-
6	pensing within such system (which may include
7	interstate shipment) shall be considered a tradi-
8	tional compounder if such pharmacy otherwise
9	meets the definition under subparagraph (A).
10	"(b) Exemptions From Certain New Drug Re-
11	QUIREMENTS.—
12	"(1) Drugs compounded by traditional
13	COMPOUNDERS.—Sections $501(a)(2)(B)$, $502(f)(1)$
14	505 (in the case of a human drug), and 512 (in the
15	case of an animal drug) shall not apply to a com-
16	pounded drug if such drug—
17	"(A) is compounded by a traditional
18	compounder that is in compliance with this sec-
19	tion; and
20	"(B) meets the requirements of this sec-
21	tion applicable to drugs compounded by tradi-
22	tional compounders.
23	"(2) Drugs compounded by compounding
24	MANUFACTURERS.—Sections 502(f)(1), 505 (in the
25	case of a human drug), and 512 (in the case of an

1	animal drug) shall not apply to a compounded drug
2	if such drug—
3	"(A) is compounded by a compounding
4	manufacturer—
5	"(i) that is not licensed as a phar-
6	macy in any State; and
7	"(ii) that is in compliance with this
8	section; and
9	"(B) meets the requirements of this sec-
10	tion applicable to drugs compounded by
11	compounding manufacturers.
12	"(c) Drugs That May Not Be Compounded.—
13	"(1) In general.—In no case may the fol-
14	lowing drugs be compounded:
15	"(A) Complex dosage forms and bio-
16	LOGICS.—A drug that is a complex dosage form
17	or biological product designated by the Sec-
18	retary pursuant to paragraph (2).
19	"(B) Marketed drugs.—A drug that is
20	a copy of a marketed drug approved under sec-
21	tion 505 or 512, including variations of such
22	drug compounded from bulk substances and a
23	drug subject to a risk evaluation and mitigation
24	strategy approved with elements to assure safe

1	use pursuant to section 505–1, as applicable,
2	except as provided in paragraph (3).
3	"(C) Drugs removed for safety and
4	EFFICACY.—A drug that appears on a list pub-
5	lished by the Secretary in the Federal Register
6	of drugs that have been withdrawn or removed
7	from the market because such drug or compo-
8	nents of such drug have been found to be un-
9	safe or not effective, subject to paragraph (4).
10	"(2) Complex dosage forms and bio-
11	LOGICS.—
12	"(A) In General.—The Secretary may
13	promulgate a regulation that designates drugs
14	or categories of drugs that are complex dosage
15	forms or biological products that may not be
16	compounded. Such regulation—
17	"(i) may include the designation of
18	drugs or categories of drugs that present
19	demonstrable difficulties for compounding,
20	such as extended release products, metered
21	dose inhalers, transdermal patches, and
22	liposomal products; and
23	"(ii) shall specify, for each drug in-
24	cluded on the list, whether the prohibition

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1	applies to the use of the drug in humans,
2	animals, or both.
3	"(B) Interim list.—
4	"(i) In general.—Before the effec-
5	tive date of the regulation promulgated
6	under subparagraph (A), the Secretary
7	may designate drugs that are complex dos-
8	age forms or biological products that can-
9	not be compounded by—
10	"(I) publishing a notice of such
11	drugs proposed for designation in the
12	Federal Register;
13	"(II) providing a period of not
14	less than 30 days for comment on the
15	notice; and
16	"(III) publishing a notice in the
17	Federal Register designating the
18	drugs that are complex dosage forms
19	and biological products that cannot be
20	compounded.
21	"(ii) Sunset.—Any notice provided
22	under clause (i) shall cease to have force or
23	effect on the date that is 5 years after the
24	date of enactment of the [insert short
25	title] or on the effective date of the final

1	regulation under subparagraph (A), which-
2	ever is earlier.
3	"(3) Exceptions regarding marketed
4	DRUGS.—A drug that is a copy of a marketed drug
5	approved under section 505 or 512, including vari-
6	ations of such drug compounded from bulk sub-
7	stances, may only be compounded if—
8	"(A) prior to beginning compounding a
9	variation of such drug, the entity compounding
10	the variation receives a prescription order for
11	an identified individual patient indicating that
12	the compounded variation produces for that pa-
13	tient a significant difference, as determined by
14	the prescribing practitioner, between the com-
15	pounded drug and the marketed drug approved
16	under section 505 or 512, as applicable;
17	"(B) in the case of a marketed drug ap-
18	proved under section 505 that is subject to a
19	risk evaluation and mitigation strategy ap-
20	proved with elements to assure safe use pursu-
21	ant to section 505-1, the entity compounding
22	the variation demonstrates to the Secretary
23	that the entity will utilize controls that are
24	comparable to the controls applicable under the

1	relevant risk evaluation and mitigation strategy;
2	or
3	"(C)(i) such marketed drug approved
4	under section 505 or 512 is included, at the
5	time of compounding a copy of such drug and
6	at the time of distribution of the compounded
7	drug—
8	"(I) in the case of a human drug, on
9	the drug shortage list under section 506E;
10	or
11	"(II) in the case of an animal drug,
12	on the Current Drug Shortages list for vet-
13	erinary products maintained on the Inter-
14	net Web site of the Food and Drug Admin-
15	istration; and
16	"(ii) the traditional compounder or the
17	compounding manufacturer notifies the Sec-
18	retary prior to the date that the compounding
19	of such a drug begins.
20	"(4) Requirement regarding drugs re-
21	MOVED FOR SAFETY OR EFFICACY.—The list pub-
22	lished by the Secretary in the Federal Register of
23	drugs that have been withdrawn or removed from
24	the market, as described in paragraph (1)(C), shall
25	specify whether a human drug on such list may, not-

1	withstanding the inclusion on such list, be com-
2	pounded for use in animals.
3	"(d) Requirements Regarding Bulk Sub-
4	STANCES APPLICABLE TO TRADITIONAL COMPOUNDERS
5	AND COMPOUNDING MANUFACTURERS.—
6	"(1) Bulk substances; human drugs.—If a
7	traditional compounder or a compounding manufac-
8	turer compounds a human drug using bulk sub-
9	stances, such compounder shall—
10	"(A) use bulk drug substances (as defined
11	in regulations of the Secretary published at sec-
12	tion 207.3(a)(4) of title 21, Code of Federal
13	Regulations (or any successor regulations))—
14	"(i) that—
15	"(I) comply with the standards of
16	an applicable United States Pharma-
17	copoeia or National Formulary mono-
18	graph, if a monograph exists, and the
19	United States Pharmacopoeia chapter
20	on pharmacy compounding;
21	"(II) if such a monograph does
22	not exist, are drug substances that
23	are components of drugs approved by
24	the Secretary; or

1	"(III) if such a monograph does
2	not exist and the drug substance is
3	not a component of a drug approved
4	by the Secretary, that appear on a list
5	developed by the Secretary through
6	regulations issued by the Secretary;
7	"(ii) that are manufactured by an es-
8	tablishment that is registered under sec-
9	tion 510 (including a foreign establishment
10	that is registered under section 510(i));
11	and
12	"(iii) that are accompanied by valid
13	certificates of analysis for each bulk drug
14	substance; and
15	"(B) use ingredients (other than bulk drug
16	substances) that comply with the standards of
17	an applicable United States Pharmacopoeia or
18	National Formulary monograph, if a mono-
19	graph exists, and the United States Pharma-
20	copoeia chapter on pharmacy compounding.
21	"(2) Bulk substances; animal drugs.—If a
22	traditional compounder or a compounding manufac-
23	turer compounds an animal drug using bulk sub-
24	stances, such compounder—

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1	"(A) shall use bulk drug substances (as de-
2	fined in regulations of the Secretary published
3	at section 207.3(a)(4) of title 21, Code of Fed-
4	eral Regulations (or any successor regulations))
5	that—
6	"(i) are manufactured by an establish-
7	ment that is registered under section 510
8	(including a foreign establishment that is
9	registered under section 510(i));
10	"(ii) are accompanied by valid certifi-
11	cates of analysis for each bulk drug sub-
12	stance; and
13	"(iii) are compounded—
14	"(I) using adequate procedures
15	and processes that ensure the safety
16	and effectiveness of the compounded
17	drug;
18	"(II) using a compounding oper-
19	ation that is commensurate with the
20	established need for compounded
21	products; and
22	"(III) following applicable State
23	laws relating to the compounding of
24	drugs for use in animals;

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1	"(B) shall use ingredients (other than bulk
2	drug substances) that comply with the stand-
3	ards of an applicable United States Pharma-
4	copoeia or National Formulary monograph, if a
5	monograph exists, and the United States Phar-
6	macopoeia chapter on pharmacy compounding;
7	"(C) in the case of a compounded animal
8	drug for use in minor species, shall use bulk
9	substances that—
10	"(i) comply with the standards of an
11	applicable United States Pharmacopoeia or
12	National Formulary monograph, if a
13	monograph exists, and the United States
14	Pharmacopoeia chapter on pharmacy
15	compounding;
16	"(ii) if such a monograph does not
17	exist, are drug substances that are compo-
18	nents of drugs approved by the Secretary;
19	or
20	"(iii) if such a monograph does not
21	exist and the drug substance is not a com-
22	ponent of a drug approved by the Sec-
23	retary, that appear on a list developed by
24	the Secretary through regulations issued
25	by the Secretary; and

1	"(D) in the case of a compounded animal
2	drug for use in major species and in food-pro-
3	ducing animals, shall use bulk substances that
4	are included on a list established by the Sec-
5	retary of bulk substances acceptable for use in
6	compounding a drug for one or both types of
7	animals.
8	"(3) Procedure.—In establishing a list of
9	designated bulk substances acceptable for use in
10	compounding a drug for use in major species or
11	food-producing animals (or both) under paragraph
12	(2)(D), the Secretary shall—
13	"(A) publish a notice of such bulk sub-
14	stances proposed for designation in the Federal
15	Register;
16	"(B) provide a period of not less than 30
17	days for comment on the notice; and
18	"(C) publish a notice in the Federal Reg-
19	ister designating the bulk substances acceptable
20	for use in compounding a drug for use in major
21	species or food-producing animals (or both).
22	"(4) WITHDRAWAL PERIODS.—The extended
23	withdrawal periods established by the Secretary pur-
24	suant to section 530.20 of title 21, Code of Federal
25	Regulations (or any successor regulations) shall

1	apply to compounded animal drugs for use in food-
2	producing animals that were compounded using bulk
3	substances.
4	"(5) Identification by secretary.—
5	"(A) IN GENERAL.—Notwithstanding the
6	existence of an applicable monograph under
7	paragraph $(1)(A)(i)(I)$ or $(2)(B)(i)(I)$, the Sec-
8	retary may identify bulk substances that the
9	Secretary determines may not be used in
10	compounding a drug.
11	"(B) PROCEDURE.—In identifying the bulk
12	substances that may not be used in
13	compounding, the Secretary shall—
14	"(i) publish a notice of such bulk sub-
15	stances proposed for identification in the
16	Federal Register;
17	"(ii) provide a period of not less than
18	30 days for comment on the notice; and
19	"(iii) publish a notice in the Federal
20	Register identifying the bulk substances
21	that may not be used in compounding a
22	drug.
23	"(e) Requirements Regarding Wholesaling
24	AND LABELING APPLICABLE TO TRADITIONAL

1	Compounders and Compounding Manufacturers.—
2	A compounded drug—
3	"(1) may not be sold by an entity other than
4	the compounding manufacturer or traditional
5	compounder that compounded the drug;
6	"(2) may not be sold to an entity other than a
7	health care entity that provides medical services
8	through licensed prescribers directly to patients, or
9	a network of such providers, except that—
10	"(A) in the case of a drug compounded by
11	a traditional compounder, the drug may be dis-
12	pensed to an individual; and
13	"(B) a compounding manufacturer may
14	transfer without profit a compounded drug to a
15	licensed pharmacy if the licensed pharmacy falls
16	under the same corporate ownership as the
17	compounding manufacturer, and the transfer of
18	such compounded drug is solely for the purpose
19	of dispensing the compounded drug to the end
20	user, who has been instructed by the pre-
21	scribing physician to self-administer such com-
22	pounded drug; and
23	"(3) in the case of a compounded drug sold to
24	a health care entity described in paragraph (2), shall
25	be labeled 'not for resale'.

1	"(f) Other Requirements Applicable to
2	Compounding Manufacturers.—
3	"(1) Licensed pharmacist oversight.—A
4	compounding manufacturer shall ensure that a phar-
5	macist licensed in the State where the compounding
6	manufacturer is located exercises direct supervision
7	over the operations of the compounding manufac-
8	turer.
9	"(2) Listing of drugs.—
10	"(A) In General.—Not less than once
11	every 6 months, a compounding manufacturer
12	shall file with the Secretary a list of—
13	"(i) the drugs compounded by such
14	compounding manufacturer during the pre-
15	vious 6-month period; and
16	"(ii) with respect to each drug listed
17	under clause (i), provide the active ingre-
18	dient, the source of such active ingredient,
19	the National Drug Code number of the
20	source drug or bulk active ingredient, the
21	strength of the active ingredient per unit,
22	the dosage form and route of administra-
23	tion, the package description, the number
24	of individual units produced, the National
25	Drug Code number of the final product,

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and other applicable requirements identi-1 2 fied by the Secretary in accordance with 3 subparagraph (B). 4 "(B) FORM.—Each list under subpara-5 graph (A) shall be prepared in such form and 6 manner as the Secretary may prescribe by regu-7 lation or guidance. "(3) Adverse event reporting.— 8 9 "(A) IN GENERAL.—If a compounding 10 manufacturer becomes aware of any serious ad-11 verse drug experience, as defined in section 12 505–1(b), such manufacturer shall report each 13 such instance to the Secretary in accordance 14 with section 314.80(c) of title 21, Code of Fed-15 eral Regulations (or any successor regulation), 16 as soon as practicable, but in no case later than 17 15 calendar days after the initial receipt of the 18 applicable information. 19 "(B) MAINTENANCE OF RECORDS.—A 20 compounding manufacturer shall maintain for a 21 period of 10 years records of all serious adverse 22 drug experiences known to the compound man-23 ufacturer in accordance with section 314.80(i)

of title 21, Code of Federal Regulations (or any

1	successor regulation), or as otherwise directed
2	by the Secretary in regulations.
3	"(4) Labeling of Drugs.—The labeling of a
4	drug compounded by a compounding manufacturer
5	shall include—
6	"(A) the statement 'This is a compounded
7	drug.' or a reasonable comparable alternative
8	statement (as specified by the Secretary) that
9	identifies the drug as a compounded drug;
10	"(B) the name, address, and phone num-
11	ber of the applicable compounding manufac-
12	turer;
13	"(C) with respect to the compounded
14	drug—
15	"(i) the lot or batch number;
16	"(ii) the established name of the
17	medication;
18	"(iii) the strength;
19	"(iv) the statement of quantity;
20	"(v) the directions for use, as appro-
21	priate;
22	"(vi) the date that the drug was com-
23	pounded;
24	"(vii) the 'beyond use' date; and
25	"(viii) storage instructions; and

1	"(D) any other information as determined
2	necessary and specified in regulations promul-
3	gated by the Secretary.
4	"(g) Compounding Manufacturer Establish-
5	MENT AND REINSPECTION FEES.—
6	"(1) Definitions.—In this subsection—
7	"(A) the term 'affiliate' has the meaning
8	given such term in section 735(11);
9	"(B) the term 'reinspection' means, with
10	respect to a compounding manufacturer, 1 or
11	more inspections conducted under section 704
12	subsequent to an inspection conducted under
13	such provision which identified noncompliance
14	materially related to an applicable requirement
15	of this Act, specifically to determine whether
16	compliance has been achieved to the Secretary's
17	satisfaction; and
18	"(C) the term 'reinspection-related costs'
19	means all expenses, including administrative ex-
20	penses, incurred in connection with—
21	"(i) arranging, conducting, and evalu-
22	ating the results of reinspections; and
23	"(ii) assessing and collecting reinspec-
24	tion fees under this subsection.

1	"(2) Establishment and reinspection
2	FEES.—For fiscal year 2015 and each subsequent
3	fiscal year, the Secretary shall, in accordance with
4	this subsection, assess and collect—
5	"(A) an establishment fee from each
6	compounding manufacturer to cover inspection-
7	related costs relating to inspections of drug
8	compounders for such year; and
9	"(B) a reinspection fee from each
10	compounding manufacturer subject to a rein-
11	spection in such fiscal year, to cover the rein-
12	spection-related costs associated with such
13	compounding manufacturer for such year.
14	"(3) Establishment fee setting.—The Sec-
15	retary shall establish the establishment fee to be col-
16	lected under this subsection for each fiscal year,
17	based on the methodology described in paragraph
18	(4) and shall publish such fee in a Federal Register
19	notice not later than 60 days before the start of
20	each such year.
21	"(4) Amount of establishment fee.—
22	"(A) In general.—Except as provided in
23	subparagraph (D), the amount of the establish-
24	ment fee under paragraph (2)(A) for a

1	compounding manufacturer in a fiscal year
2	shall be equal to the sum of—
3	(i)(I) \$15,000 per drug establish-
4	ment owned or operated by the
5	compounding manufacturer, multiplied by
6	"(II) the inflation adjustment factor
7	described in subparagraph (B); plus
8	"(ii) the small business adjustment
9	factor described in subparagraph (C).
10	"(B) Inflation adjustment factor.—
11	"(i) In general.—For fiscal year
12	2015 and subsequent fiscal years, the reve-
13	nues established in subparagraph (A) shall
14	be adjusted by the Secretary by notice,
15	published in the Federal Register, for a
16	fiscal year by the amount equal to the sum
17	of—
18	``(I) one;
19	"(II) the average annual percent
20	change in the cost, per full-time equiv-
21	alent position of the Food and Drug
22	Administration, of all personnel com-
23	pensation and benefits paid with re-
24	spect to such positions for the first 3
25	years of the preceding 4 fiscal years,

1	multiplied by the proportion of per-
2	sonnel compensation and benefits
3	costs to total costs of conducting in-
4	spections of drug compounders for the
5	first 3 years of the preceding 4 fisca
6	years, and
7	"(III) the average annual percent
8	change that occurred in the Consumer
9	Price Index for urban consumers
10	(Washington-Baltimore, DC-MD-VA-
11	WV; Not Seasonally Adjusted; Al
12	items; Annual Index) for the first 3
13	years of the preceding 4 years of
14	available data multiplied by the pro-
15	portion of all costs other than per-
16	sonnel compensation and benefits
17	costs to total costs of conducting in-
18	spections of drug compounders for the
19	first 3 years of the preceding 4 fisca
20	years.
21	"(ii) COMPOUNDED BASIS.—The ad-
22	justment made each fiscal year under
23	clause (i) shall be added on a compounded
24	basis to the sum of all adjustments made

1	each fiscal year after fiscal year 2014
2	under clause (i).
3	"(C) SMALL BUSINESS ADJUSTMENT FAC-
4	TOR.—The small business adjustment factor de-
5	scribed in subparagraph (A)(ii) shall be an
6	amount established by the Secretary for each
7	fiscal year based on the Secretary's estimate
8	of—
9	"(i) the number of small businesses
10	that will pay a reduced establishment fee
11	for such fiscal year; and
12	"(ii) the adjustment to the establish-
13	ment fee necessary to achieve total fees
14	equaling the total fees that the Secretary
15	would have collected if no entity qualified
16	for the small business exception in sub-
17	paragraph (D).
18	"(D) Exception for small busi-
19	NESSES.—In the case of a compounding manu-
20	facturer that employs 25 or less employees, in-
21	cluding employees of an affiliate, the amount of
22	the establishment fee under paragraph (2)(A)
23	for a fiscal year shall be equal to $\frac{1}{3}$ of the
24	amount calculated under subparagraph (A)(i) in
25	such fiscal year.

1 "(E) CREDITING OF FEES.—In estab-2 lishing the small business adjustment factor 3 under this paragraph for a fiscal year, the Sec-4 retary shall provide for the crediting of fees 5 from the previous year to the next year if the 6 Secretary overestimated the amount of the 7 small business adjustment factor for such pre-8 vious fiscal year, and consider the need to ac-9 count for any adjustment of fees and such other 10 factors as the Secretary determines appropriate. 11 "(5) Amount of reinspection fee.—The 12 amount of the reinspection fee under paragraph 13 (2)(B) for a compounding manufacturer in a fiscal 14 vear shall be the amount that is 100 percent of the 15 reinspection-related costs (including by type or level 16 of reinspection activity, as the Secretary determines 17 applicable) applicable to such compounding manu-18 facturer in such year. 19 "(6) Use of fees.—The Secretary shall make 20 all of the fees collected pursuant to subparagraph 21 (A) and (B) of paragraph (2) available solely to pay 22 for the costs referred to in such subparagraph (A) 23 or (B) of paragraph (2), respectively. "(7) SUPPLEMENT NOT SUPPLANT.—Funds re-24 25 ceived by the Secretary pursuant to this section shall

1 be used to supplement and not supplant any other 2 Federal funds available to carry out the activities de-3 scribed in this section. "(8) Crediting and availability of fees.— 4 5 Fees authorized under this subsection shall be col-6 lected and available for obligation only to the extent 7 and in the amount provided in appropriations Acts. 8 Such fees are authorized to remain available until 9 expended. Such sums as may be necessary may be 10 transferred from the Food and Drug Administration 11 salaries and expenses account without fiscal year 12 limitation to such appropriation account for salaries 13 and expenses with such fiscal year limitation. The 14 sums transferred shall be available solely for the 15 purpose of paying the operating expenses of the 16 Food and Drug Administration employees and con-17 tractors performing activities with respect to the 18 oversight of compounded drugs. 19 "(9) Collection of fees.— 20 "(A) ESTABLISHMENT FEE.—A 21 compounding manufacturer shall remit the es-22 tablishment fee due under this subsection in a 23 fiscal year when submitting a registration pur-24 suant to section 510(g) for such fiscal year. A 25 compounding manufacturer shall not be consid-

1 ered registered under section 510 in a fiscal 2 year until the date that the compounding man-3 ufacturer remits the establishment fee under 4 this subsection for such fiscal year. 5 "(B) Reinspection fee.—The Secretary 6 shall specify in the Federal Register notice de-7 scribed in paragraph (3) the time and manner 8 in which reinspection fees assessed under this 9 subsection shall be collected. Such a fee shall be 10 collected after the Secretary has conducted a 11 reinspection of the compounding manufacturer 12 involved. 13 "(C) COLLECTION OF UNPAID FEES.—In 14 any case where the Secretary does not receive 15 payment of a fee assessed under this subsection 16 within 30 days after it is due, such fee shall be 17 treated as a claim of the United States Govern-18 ment subject to provisions of subchapter II of 19 chapter 37 of title 31, United States Code. "(10) Annual Report to Congress.—Not 20 21 later than 120 days after each fiscal year in which 22 fees are assessed under this subsection, the Sec-23 retary shall submit a report to the Committee on 24 Health Education Labor and Pensions of the Senate 25 and the Committee on Energy and Commerce of the

1 House of Representatives, to include a description of 2 fees assessed and collected for each year, a summary 3 description of entities paying the fees, and the num-4 ber of inspections and reinspections of such entities 5 performed each year. "(11) AUTHORIZATION OF APPROPRIATIONS.— 6 7 For fiscal year 2015 and each subsequent fiscal 8 year, there is authorized to be appropriated for fees 9 under this subsection an amount equal to the total 10 revenue amount estimated by the Secretary to be 11 collected pursuant to paragraphs (4) and (5) for the 12 fiscal year, as adjusted or otherwise affected under 13 the other provisions of this subsection. 14 "(h) Action by Secretary Regarding Com-15 PLAINTS FROM STATE BOARDS OF PHARMACY.—The Secretary shall encourage direct communication between 16 17 States regarding traditional compounders. If the Secretary 18 receives a complaint from a State Board of Pharmacy, in 19 a manner specified by the Secretary, about compounded 20 drugs produced by an identified traditional compounder 21 licensed as a pharmacy in another identified State, the 22 Secretary shall notify the identified State in which the tra-23 ditional compounder is licensed of the complaint within 15 days. This obligation of the Secretary does not expand the

- 1 Secretary's authority over or responsibility for the tradi-
- 2 tional compounder that is the subject of the complaint.
- 3 "(i) Prescription Order Reference.—For pur-
- 4 poses of this section, reference to a prescription order for
- 5 an identified individual patient includes, in the case of ani-
- 6 mal drugs, a prescription order for a specific herd or flock
- 7 of animals.
- 8 "(j) APPLICATION.—This section shall not apply to
- 9 compounded positron emission tomography drugs.".
- 10 (c) Prohibited Act.—Section 301 (21 U.S.C. 331)
- 11 is amended by adding at the end the following:
- 12 "(ccc) The resale of a compounded drug that is la-
- 13 beled 'not for resale' as required by section 503A.".
- 14 (d) REPORT BY GAO.—Not later than November 1,
- 15 2016, the Comptroller General of the United States shall
- 16 conduct study and submit to Congress a report regarding
- 17 the impact of this Act (and the amendments made by this
- 18 Act) on the safety of animal drug compounding and the
- 19 availability of safe and effective drugs for animals.
- 20 SEC. 3. OTHER REQUIREMENTS RELATING TO
- 21 COMPOUNDING MANUFACTURERS.
- 22 (a) Labeling.—Section 502 (21 U.S.C. 352) is
- 23 amended by adding at the end the following:

1 "(bb) If it is a compounded drug and the labeling 2 does not include the information as required by sub-3 sections (e)(3) and (f)(4) of section 503A, as applicable.". 4 (b) Application of Manufacturer Registra-5 TION REQUIREMENTS TO COMPOUNDING MANUFACTUR-ERS.—Section 510(g) (21 U.S.C. 360(g)) is amended— 6 7 (1) by redesignating paragraphs (1) through 8 (5) as paragraphs (A) through (E); 9 (2) by striking "(g) The foregoing" and insert-10 ing "(g)(1) The foregoing"; and 11 (3) by inserting after subparagraph (E), as so 12 redesignated, the following: 13 "(2)(A) The exemption in paragraph (1)(A) does not apply with respect to compounding manufacturers. The 14 15 Secretary shall establish procedures to provide for the registration under this section of compounding manufactur-16 17 ers, which shall include the payment of the establishment fee under section 503A(f). As part of these procedures, 18 the Secretary shall establish a timeline for registration for 19 20 the first year following the date of enactment of the Lin-21 sert short title. In no case may registration be required 22 until at least 60 days following publication of the timeline 23 in the Federal Register, notwithstanding subsection 24 (b)(1).

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1 "(B) In this subsection, the term compounding man-2 ufacturer' has the meaning given such term in section 3 503A.". 4 (c) Application of Inspection Requirements to 5 Compounding Manufacturers.—Section 704(a)(2)(21 U.S.C. 374(a)(2)) is amended by adding at the end 6 7 the following flush text: 8 "The exemption in subparagraph (A) does not apply with respect to compounding manufacturers (as such term is 10 defined in section 503A).". SEC. 4. IMPLEMENTATION. 12 In promulgating any regulations to implement this 13 Act (and the amendments made by this Act), the Sec-14 retary of Health and Human Services shall— 15 (1) issue a notice of proposed rulemaking that 16 includes the proposed regulation; 17 (2) provide a period of not less than 60 days 18 for comments on the proposed regulation; and 19 (3) publish the final regulation not more than 20 18 months following publication of the proposed rule

and not less than 30 days before the effective date

of such final regulation.