To amend the Public Health Service Act to establish a pathway for the licensure of biosimilar biological products, to promote innovation in the life sciences, and for other purposes.

IN THE SENATE OF THE UNITED STATES

Mr.	Kennedy	(for himse	elf, Mr. H	ATCH, Mrs	. Clint	ON,	and Mr.	Enzi)	intro-
	duced the	following	bill; which	h was reac	l twice	and	referred	to the	Com-
	mittee on								

A BILL

- To amend the Public Health Service Act to establish a pathway for the licensure of biosimilar biological products, to promote innovation in the life sciences, 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 "Biologics Price Com-
 - 5 petition and Innovation Act of 2007".

1	SEC. 2. APPROVAL PATHWAY FOR BIOSIMILAR BIOLOGICAL
2	PRODUCTS.
3	(a) Licensure of Biological Products as Bio-
4	SIMILAR OR INTERCHANGEABLE.—Section 351 of the
5	Public Health Service Act (42 U.S.C. 262) is amended—
6	(1) in subsection $(a)(1)(A)$, by inserting "under
7	this subsection or subsection (k)" after "biologics li-
8	cense"; and
9	(2) by adding at the end the following:
10	"(k) Licensure of Biological Products as Bio-
11	SIMILAR OR INTERCHANGEABLE.—
12	"(1) In general.—Any person may submit an
13	application for licensure of a biological product
14	under this subsection.
15	"(2) Content.—
16	"(A) In general.—
17	"(i) Required information.—An
18	application submitted under this subsection
19	shall include information demonstrating
20	that—
21	"(I) the biological product is bio-
22	similar to a reference product based
23	upon data derived from—
24	"(aa) analytical studies that
25	demonstrate that the biological
26	product is highly similar to the

1	reference product notwith-
2	standing minor differences in
3	clinically inactive components;
4	"(bb) animal studies; and
5	"(ce) a clinical study or
6	studies (including the assessment
7	of immunogenicity and phar-
8	macokinetics
9	pharmacodynamics) that are—
10	"(AA) sufficient to
11	demonstrate safety, purity,
12	and potency in 1 or more
13	appropriate conditions of use
14	for which the reference
15	product is licensed and in-
16	tended to be used and for
17	which licensure is sought for
18	the biological product; and
19	"(BB) designed to
20	avoid needlessly duplicative
21	or unethical clinical testing;
22	"(II) the biological product and
23	reference product utilize the same
24	mechanism or mechanisms of action
25	for the condition or conditions of use

1	prescribed, recommended, or sug-
2	gested in the proposed labeling, but
3	only to the extent the mechanism or
4	mechanisms of action are known for
5	the reference product;
6	"(III) the condition or conditions
7	of use prescribed, recommended, or
8	suggested in the labeling proposed for
9	the biological product have been pre-
10	viously approved for the reference
11	product;
12	"(IV) the route of administra-
13	tion, the dosage form, and the
14	strength of the biological product are
15	the same as those of the reference
16	product; and
17	"(V) the facility in which the bio-
18	logical product is manufactured, proc-
19	essed, packed, or held meets stand-
20	ards designed to assure that the bio-
21	logical product continues to be safe,
22	pure, and potent.
23	"(ii) Determination by sec-
24	RETARY.—The Secretary may determine,
25	in the Secretary's discretion, that an ele-

1	ment described in clause (i)(I) is unneces-
2	sary in an application submitted under this
3	subsection.
4	"(iii) Additional information.—
5	An application submitted under this sub-
6	section may include—
7	"(I) at the applicant's option,
8	publicly-available information regard-
9	ing the Secretary's previous deter-
10	mination that the reference product is
11	safe, pure, and potent; and
12	"(II) any additional information
13	in support of the application, includ-
14	ing publicly-available information with
15	respect to the reference product or an-
16	other biological product.
17	"(B) Interchangeability.—An applica-
18	tion (or a supplement to an application) sub-
19	mitted under this subsection may include infor-
20	mation demonstrating that the biological prod-
21	uct is interchangeable with the reference prod-
22	uct.
23	"(3) Evaluation by secretary.—Upon re-
24	view of an application (or a supplement to an appli-
25	cation) submitted under this subsection, the Sec-

1	retary shall license the biological product under this
2	subsection if the Secretary determines that the infor-
3	mation submitted in the application (or the supple-
4	ment) is sufficient to show that the biological prod-
5	uct—
6	"(A) is biosimilar to the reference product
7	or
8	"(B) is interchangeable with the reference
9	product.
10	"(4) SAFETY STANDARDS FOR DETERMINING
11	INTERCHANGEABILITY.—Upon review of an applica-
12	tion submitted under this subsection or any supple-
13	ment to such application, the Secretary shall deter-
14	mine the biological product to be interchangeable
15	with the reference product if the Secretary deter-
16	mines that the information submitted in the applica-
17	tion (or a supplement to such application) is suffi-
18	cient to show that—
19	"(A) the biological product—
20	"(i) is biosimilar to the reference
21	product; and
22	"(ii) can be expected to produce the
23	same clinical result as the reference prod-
24	uct in any given patient; and

25

1	"(B) for a biological product that is ad-
2	ministered more than once to an individual, the
3	risk in terms of safety or diminished efficacy of
4	alternating or switching between use of the bio-
5	logical product and the reference product is not
6	greater than the risk of using the reference
7	product without such alternation or switch.
8	"(5) General rules.—
9	"(A) ONE REFERENCE PRODUCT PER AP-
10	PLICATION.—A biological product, in an appli-
11	cation submitted under this subsection, may not
12	be evaluated against more than 1 reference
13	product.
14	"(B) Review.—An application submitted
15	under this subsection shall be reviewed by the
16	division within the Food and Drug Administra-
17	tion that is responsible for the review and ap-
18	proval of the application under which the ref-
19	erence product is licensed.
20	"(C) RISK EVALUATION AND MITIGATION
21	STRATEGIES.—The authority of the Secretary
22	with respect to risk evaluation and mitigation
23	strategies under the Federal Food, Drug, and

Cosmetic Act shall apply to biological products

licensed under this subsection in the same man-

1	ner as such authority applies to biological prod-
2	ucts licensed under subsection (a).
3	"(6) Exclusivity for first interchange-
4	ABLE BIOLOGICAL PRODUCT.—Upon review of an
5	application submitted under this subsection relying
6	on the same reference product for which a prior bio-
7	logical product has received a determination of inter-
8	changeability for any condition of use, the Secretary
9	shall not make a determination under paragraph (4)
10	that the second or subsequent biological product is
11	interchangeable for any condition of use until the
12	earlier of—
13	"(A) 1 year after the first commercial
14	marketing of the first interchangeable bio-
15	similar biological product to be approved as
16	interchangeable for that reference product;
17	"(B) 18 months after—
18	"(i) a final court decision on all pat-
19	ents in suit in an action instituted under
20	subsection (l)(6) against the applicant that
21	submitted the application for the first ap-
22	proved interchangeable biosimilar biological
23	product; or
24	"(ii) the dismissal with or without
25	prejudice of an action instituted under sub-

1	section (l)(6) against the applicant that
2	submitted the application for the first ap-
3	proved interchangeable biosimilar biological
4	product; or
5	"(C)(i) 42 months after approval of the
6	first interchangeable biosimilar biological prod-
7	uct if the applicant that submitted such appli-
8	cation has been sued under subsection (l)(6)
9	and such litigation is still ongoing within such
10	36-month period; or
11	"(ii) 18 months after approval of the first
12	interchangeable biosimilar biological product if
13	the applicant that submitted such application
14	has not been sued under subsection (l)(6).
15	For purposes of this paragraph, the term 'final court
16	decision' means a final decision of a court from
17	which no appeal (other than a petition to the United
18	States Supreme Court for a writ of certiorari) has
19	been or can be taken.
20	"(7) Exclusivity for reference prod-
21	UCT.—
22	"(A) Effective date of biosimilar ap-
23	PLICATION APPROVAL.—Approval of an applica-
24	tion under this subsection may not be made ef-
25	fective by the Secretary until the date that is

1	12 years after the date on which the reference
2	product was first licensed under subsection (a).
3	"(B) FILING PERIOD.—An application
4	under this subsection may not be submitted to
5	the Secretary until the date that is 4 years
6	after the date on which the reference product
7	was first licensed under subsection (a).
8	"(8) Guidance documents.—
9	"(A) In General.—The Secretary may,
10	after opportunity for public comment, issue
11	guidance in accordance, except as provided in
12	subparagraph (B)(i), with section 701(h) of the
13	Federal Food, Drug, and Cosmetic Act with re-
14	spect to the process for the submission of appli-
15	cations for, and licensure of, a biological prod-
16	uct under this subsection. Any such guidance
17	may be general or specific.
18	"(B) Public comment.—
19	"(i) In General.—The Secretary
20	shall provide the public an opportunity to
21	comment on any proposed guidance issued
22	under subparagraph (A) before issuing
23	final guidance.
24	"(ii) Input regarding most valu-
25	ABLE GUIDANCE.—The Secretary shall es-

1	tablish a process through which the public
2	may provide the Secretary with input re-
3	garding priorities for issuing guidance.
4	"(C) No requirement for application
5	CONSIDERATION.—The issuance (or non-
6	issuance) of guidance under subparagraph (A)
7	shall not preclude the review of, or action on,
8	an application submitted under this subsection.
9	"(D) REQUIREMENT FOR PRODUCT CLASS-
10	SPECIFIC GUIDANCE.—If the Secretary issues
11	product class-specific guidance under subpara-
12	graph (A), such guidance shall include a de-
13	scription of—
14	"(i) the criteria that the Secretary will
15	use to determine whether a biological prod-
16	uct is highly similar to a reference product
17	in such product class; and
18	"(ii) the criteria, if available, that the
19	Secretary will use to determine whether a
20	biological product meets the standards de-
21	scribed in paragraph (4).
22	"(E) CERTAIN PRODUCT CLASSES.—
23	"(i) Guidance.—The Secretary may
24	indicate in a guidance document that the
25	science and experience, as of the date of

1	such guidance, with respect to a product or
2	product class (not including any recom-
3	binant protein) does not allow approval of
4	an application for a license as provided
5	under this subsection for such product or
6	product class.
7	"(ii) Modification or reversal.—
8	The Secretary may issue a subsequent
9	guidance document under subparagraph
10	(A) to modify or reverse a guidance docu-
11	ment under clause (i).
12	"(iii) No effect on ability to
13	DENY LICENSE.—Clause (i) shall not be
14	construed to require the Secretary to ap-
15	prove a product with respect to which the
16	Secretary has not indicated in a guidance
17	document that the science and experience
18	as described in clause (i), does not allow
19	approval of such an application.
20	"(l) Patents.—
21	"(1) Confidential access to subsection
22	(k) APPLICATION.—
23	"(A) APPLICATION OF PARAGRAPH.—Un-
24	less otherwise agreed to by a person that sub-
25	mits an application under subsection (k) (re-

ferred to in this subsection as the 'subsection (k) applicant') and the sponsor of the application for the reference product (referred to in this paragraph as the 'reference product sponsor'), the provisions of this paragraph shall apply to the exchange of information described in this subsection.

"(B) In General.—

"(i) Provision of confidential information,"

FORMATION.—When a subsection (k) applicant submits an application under subsection (k), such applicant shall provide to the persons described in clause (ii), subject to the terms of this paragraph, confidential access to the information required to be produced pursuant to paragraph (2) and any other information that the subsection (k) applicant determines, in its sole discretion, to be appropriate (referred to in this subsection as the 'confidential information').

"(ii) RECIPIENTS OF INFORMATION.—
The persons described in this clause are
the following:

1	"(I) Outside counsel.—One or
2	more attorneys designated by the ref-
3	erence product sponsor who are em-
4	ployees of an entity other than the
5	reference product sponsor (referred to
6	in this paragraph as the 'outside
7	counsel'), provided that such attor-
8	neys do not engage, formally or infor-
9	mally, in patent prosecution relevant
10	or related to the reference product.
11	"(II) In-house counsel.—One
12	attorney that represents the reference
13	product sponsor who is an employee
14	of the reference product sponsor, pro-
15	vided that such attorney does not en-
16	gage, formally or informally, in patent
17	prosecution relevant or related to the
18	reference product.
19	"(C) Limitation on disclosure.—No
20	person that receives confidential information
21	pursuant to subparagraph (B) shall disclose
22	any confidential information to any other per-
23	son or entity, including the reference product
24	sponsor employees, outside scientific consult-
25	ants, or other outside counsel retained by the

reference product sponsor, without the prior written consent of the subsection (k) applicant, which shall not be unreasonably withheld.

"(D) USE OF CONFIDENTIAL INFORMATION.—Confidential information shall be used
for the sole and exclusive purpose of determining, with respect to each patent assigned to
or exclusively licensed by the reference product
sponsor, whether a claim of patent infringement
could reasonably be asserted if the subsection
(k) applicant engaged in the manufacture, use,
offering for sale, sale, or importation into the
United States of the biological product that is
the subject of the application under subsection
(k).

"(E) Ownership of confidential information disformation.—The confidential information disclosed under this paragraph is, and shall remain, the property of the subsection (k) applicant. By providing the confidential information pursuant to this paragraph, the subsection (k) applicant does not provide the reference product sponsor or the outside counsel any interest in or license to use the confidential information, for

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purposes other than those specified in subparagraph (D).

> "(F) EFFECT OFINFRINGEMENT AC-TION.—In the event that the reference product sponsor files a patent infringement suit, the use of confidential information shall continue to be governed by the terms of this paragraph until such time as a court enters a protective order regarding the information. Upon entry of such order, the subsection (k) applicant may redesignate confidential information in accordance with the terms of that order. No confidential information shall be included in any publiclyavailable complaint or other pleading. In the event that the reference product sponsor does not file an infringement action by the date specified in paragraph (6), the reference product sponsor shall return or destroy all confidential information received under this paragraph, provided that if the reference product sponsor opts to destroy such information, it will confirm destruction in writing to the subsection (k) applicant.

> "(G) Rule of construction.—Nothing in this paragraph shall be construed—

1	"(i) as an admission by the subsection
2	(k) applicant regarding the validity, en-
3	forceability, or infringement of any patent;
4	or
5	"(ii) an agreement or admission by
6	the subsection (k) applicant with respect to
7	the competency, relevance, or materiality
8	of any confidential information.
9	"(H) Effect of violation.—The disclo-
10	sure of any confidential information in violation
11	of this paragraph shall be deemed to cause the
12	subsection (k) applicant to suffer irreparable
13	harm for which there is no adequate legal rem-
14	edy and the court shall consider immediate in-
15	junctive relief to be an appropriate and nec-
16	essary remedy for any violation or threatened
17	violation of this paragraph.
18	"(2) Subsection (k) application informa-
19	TION.—Not later than 20 days after the Secretary
20	notifies the subsection (k) applicant that the applica-
21	tion has been accepted for review, the subsection (k)
22	applicant—
23	"(A) shall provide to the reference product
24	sponsor a copy of the application submitted to
25	the Secretary under subsection (k), and such

1	other information that describes the process or
2	processes used to manufacture the biological
3	product that is the subject of such application;
4	and
5	"(B) may provide to the reference product
6	sponsor additional information requested by or
7	on behalf of the reference product sponsor.
8	"(3) List and description of patents.—
9	"(A) List by reference product spon-
10	SOR.—Not later than 60 days after the receipt
11	of the application and information under para-
12	graph (2), the reference product sponsor shall
13	provide to the subsection (k) applicant—
14	"(i) a list of patents for which the ref-
15	erence product sponsor believes a claim of
16	patent infringement could reasonably be
17	asserted by the reference product sponsor
18	if a person not licensed by the reference
19	product sponsor engaged in the making,
20	using, offering to sell, selling, or importing
21	into the United States of the biological
22	product that is the subject of the sub-
23	section (k) application; and
24	"(ii) an identification of the patents
25	on such list that the reference product

1	sponsor would be prepared to license to the
2	subsection (k) applicant.
3	"(B) List and description by sub-
4	SECTION (k) APPLICANT.—Not later than 60
5	days after receipt of the list under subpara-
6	graph (A), the subsection (k) applicant—
7	"(i) may provide to the reference
8	product sponsor a list of patents to which
9	the subsection (k) applicant believes a
10	claim of patent infringement could reason-
11	ably be asserted by the reference product
12	sponsor if a person not licensed by the ref-
13	erence product sponsor engaged in the
14	making, using, offering to sell, selling, or
15	importing into the United States of the bi-
16	ological product that is the subject of the
17	subsection (k) application;
18	"(ii) shall provide to the reference
19	product sponsor, with respect to each pat-
20	ent listed by the reference product sponsor
21	under subparagraph (A) or listed by the
22	subsection (k) applicant under clause (i)—
23	"(I) a detailed statement that de-
24	scribes, on a claim by claim basis, the
25	factual and legal basis of the opinion

1	of the subsection (k) applicant that
2	such patent is invalid, unenforceable
3	or will not be infringed by the com-
4	mercial marketing of the biological
5	product that is the subject of the sub-
6	section (k) application; or
7	"(II) a statement that the sub-
8	section (k) applicant does not intend
9	to begin commercial marketing of the
10	biological product before the date that
11	such patent expires; and
12	"(iii) shall provide to the reference
13	product sponsor a response regarding each
14	patent identified by the reference product
15	sponsor under subparagraph (A)(ii).
16	"(C) Description by reference prod-
17	UCT SPONSOR.—Not later than 60 days after
18	receipt of the list and statement under subpara-
19	graph (B), the reference product sponsor shall
20	provide to the subsection (k) applicant a de-
21	tailed statement that describes, with respect to
22	each patent described in subparagraph
23	(B)(ii)(I), on a claim by claim basis, the factual
24	and legal basis of the opinion of the reference
25	product sponsor that such patent will be in-

fringed by the commercial marketing of the biological product that is the subject of the subsection (k) application and a response to the statement concerning validity and enforceability provided under subparagraph (B)(ii)(I).

"(4) Patent resolution negotiations.—

"(A) IN GENERAL.—After receipt by the subsection (k) applicant of the statement under paragraph (3)(C), the reference product sponsor and the subsection (k) applicant shall engage in good faith negotiations to agree on which, if any, patents listed under paragraph (3) by the subsection (k) applicant or the reference product sponsor shall be the subject of an action for patent infringement under paragraph (6).

"(B) Failure to reach agreement.—

If, within 15 days of beginning negotiations under subparagraph (A), the subsection (k) applicant and the reference product sponsor fail to agree on a final and complete list of which, if any, patents listed under paragraph (3) by the subsection (k) applicant or the reference product sponsor shall be the subject of an action for patent infringement under paragraph (6), the

1	provisions of paragraph (5) shall apply to the
2	parties.
3	"(5) Patent resolution if no agree-
4	MENT.—
5	"(A) Number of Patents.—The sub-
6	section (k) applicant shall notify the reference
7	product sponsor of the number of patents that
8	such applicant will provide to the reference
9	product sponsor under subparagraph $(B)(i)(I)$
10	"(B) Exchange of patent lists.—
11	"(i) In general.—On a date agreed
12	to by the subsection (k) applicant and the
13	reference product sponsor, but in no case
14	later than 5 days after the subsection (k)
15	application notifies the reference product
16	sponsor under subparagraph (A), the sub-
17	section (k) applicant and the reference
18	product sponsor shall simultaneously ex-
19	change—
20	"(I) the list of patents that the
21	subsection (k) applicant believes
22	should be the subject of an action for
23	patent infringement under paragraph
24	(6); and

1	"(II) the list of patents, in ac-
2	cordance with clause (ii), that the ref-
3	erence product sponsor believes should
4	be the subject of an action for patent
5	infringement under paragraph (6).
6	"(ii) Number of patents listed by
7	REFERENCE PRODUCT SPONSOR.—
8	"(I) In general.—Subject to
9	subclause (II), the number of patents
10	listed by the reference product spon-
11	sor under clause (i)(II) may not ex-
12	ceed the number of patents listed by
13	the subsection (k) applicant under
14	clause $(i)(I)$.
15	"(II) Exception.—If a sub-
16	section (k) applicant does not list any
17	patent under clause (i)(I), the ref-
18	erence product sponsor may list 1 pat-
19	ent under clause $(i)(II)$.
20	"(6) Immediate patent infringement ac-
21	TION.—
22	"(A) ACTION IF AGREEMENT ON PATENT
23	LIST.—If the subsection (k) applicant and the
24	reference product sponsor agree on patents as
25	described in paragraph (4), not later than 30

1	days after such agreement, the reference prod-
2	uct sponsor shall bring an action for patent in-
3	fringement with respect to each such patent.
4	"(B) ACTION IF NO AGREEMENT ON PAT-
5	ENT LIST.—If the provisions of paragraph (5)
6	apply to the parties as described in paragraph
7	(4)(B), not later than 30 days after the ex-
8	change of lists under paragraph (5)(B), the ref-
9	erence product sponsor shall bring an action for
10	patent infringement with respect to each patent
11	that is included on such lists.
12	"(C) Notification and publication of
13	COMPLAINT.—
14	"(i) Notification to secretary.—
15	Not later than 30 days after a complaint
16	is served to a subsection (k) applicant in
17	an action for patent infringement described
18	under this paragraph, the subsection (k)
19	applicant shall provide the Secretary with
20	notice and a copy of such complaint.
21	"(ii) Publication by secretary.—
22	The Secretary shall publish in the Federal
23	Register notice of a complaint received
24	under clause (i).

1	"(7) Newly issued or licensed patents.—
2	In the case of a patent that—
3	"(A) is issued to, or exclusively licensed by,
4	the reference product sponsor after the date
5	that the reference product sponsor provided the
6	list to the subsection (k) applicant under para-
7	graph $(3)(A)$; and
8	"(B) the reference product sponsor reason-
9	ably believes that, due to the issuance of such
10	patent, a claim of patent infringement could
11	reasonably be asserted by the reference product
12	sponsor if a person not licensed by the ref-
13	erence product sponsor engaged in the making,
14	using, offering to sell, selling, or importing into
15	the United States of the biological product that
16	is the subject of the subsection (k) application,
17	not later than 30 days after such issuance or licens-
18	ing, the reference product sponsor shall provide to
19	the subsection (k) applicant a supplement to the list
20	provided by the reference product sponsor under
21	paragraph (3)(A) that includes such patent, not
22	later than 30 days after such supplement is pro-
23	vided, the subsection (k) applicant shall provide a
24	statement to the reference product sponsor in ac-

1	cordance with paragraph (3)(B), and such patent
2	shall be subject to paragraph (8).
3	"(8) Notice of commercial marketing and
4	PRELIMINARY INJUNCTION.—
5	"(A) NOTICE OF COMMERCIAL MAR-
6	KETING.—The subsection (k) applicant shall
7	provide notice to the reference product sponsor
8	not later than 180 days before the date of the
9	first commercial marketing of the biological
10	product licensed under subsection (k).
11	"(B) Preliminary injunction.—After
12	receiving the notice under subparagraph (A)
13	and before such date of the first commercial
14	marketing of such biological product, the ref-
15	erence product sponsor may seek a preliminary
16	injunction prohibiting the subsection (k) appli-
17	cant from engaging in the commercial manufac-
18	ture or sale of such biological product until the
19	court decides the issue of patent validity, en-
20	forcement, and infringement with respect to any
21	patent that is—
22	"(i) included in the list provided by
23	the reference product sponsor under para-
24	graph (3)(A) or in the list provided by the

1	subsection (k) applicant under paragraph
2	(3)(B); and
3	"(ii) not included, as applicable, on—
4	"(I) the list of patents described
5	in paragraph (4); or
6	"(II) the lists of patents de-
7	scribed in paragraph (5)(B).
8	"(C) REASONABLE COOPERATION.—If the
9	reference product sponsor has sought a prelimi-
10	nary injunction under subparagraph (B), the
11	reference product sponsor and the subsection
12	(k) applicant shall reasonably cooperate to ex-
13	pedite such further discovery as is needed in
14	connection with the preliminary injunction mo-
15	tion.
16	"(9) Limitation on declaratory judgment
17	ACTION.—
18	"(A) Subsection (k) application pro-
19	VIDED.—If a subsection (k) applicant provides
20	the application and information required under
21	paragraph (2)(A), neither the reference product
22	sponsor nor the subsection (k) applicant may,
23	prior to the date notice is received under para-
24	graph (8)(A), bring any action under section
25	2201 of title 28, United States Code, for a dec-

laration of infringement, validity, or enforceability of any patent that is described in clauses

(i) and (ii) of paragraph (8)(B).

"(B) Subsequent failure to act by

Subsection (k) Applicant.—If a subsection

SUBSECTION (k) APPLICANT.—If a subsection (k) applicant fails to complete an action required of the subsection (k) applicant under paragraph (3)(B)(ii), paragraph (5), paragraph (6)(C)(i), paragraph (7), or paragraph (8)(A), the reference product sponsor, but not the subsection (k) applicant, may bring an action under section 2201 of title 28, United States Code, for a declaration of infringement, validity, or enforceability of any patent included in the list described in paragraph (3)(A), including as provided under paragraph (7).

"(C) Subsection (k) application not provide the application and information required under paragraph (2)(A), the reference product sponsor, but not the subsection (k) applicant, may bring an action under section 2201 of title 28, United States Code, for a declaration of infringement, validity, or enforceability

1	of any patent that claims the biological product
2	or a use of the biological product.".
3	(b) Definitions.—Section 351(i) of the Public
4	Health Service Act (42 U.S.C. 262(i)) is amended—
5	(1) by striking "In this section, the term bio-
6	logical product' means" and inserting the following:
7	"In this section:
8	"(1) The term 'biological product' means";
9	(2) in paragraph (1), as so designated, by in-
10	serting "protein (except any chemically synthesized
11	polypeptide)," after "allergenic product,"; and
12	(3) by adding at the end the following:
13	"(2) The term 'biosimilar' or 'biosimilarity', in
14	reference to a biological product that is the subject
15	of an application under subsection (k), means there
16	are no clinically meaningful differences between the
17	biological product and the reference product in
18	terms of the safety, purity, and potency of the prod-
19	uct.
20	"(3) The term 'interchangeable' or 'inter-
21	changeability', in reference to a biological product
22	that is the subject of an application under sub-
23	section (k), means that the biological product may
24	be substituted for the reference product without the

1	intervention of the health care provider who pre-
2	scribed the reference product.
3	"(4) The term 'reference product' means the
4	single biological product licensed under subsection
5	(a) against which a biological product is evaluated in
6	an application submitted under subsection (k).".
7	(c) Conforming Amendments Relating to Pat-
8	ENTS.—
9	(1) Patents.—Section 271(e) of title 35,
10	United States Code, is amended—
11	(A) in paragraph (2)—
12	(i) in subparagraph (A), by striking
13	"or" at the end;
14	(ii) in subparagraph (B), by adding
15	"or" at the end; and
16	(iii) by inserting after subparagraph
17	(B) the following:
18	"(C)(i) with respect to a patent that is identi-
19	fied in the list of patents described in section
20	351(l)(3) of the Public Health Service Act (including
21	as provided under section 351(l)(7) of such Act), an
22	application seeking approval of a biological product,
23	or
24	"(ii) if the applicant for the application fails to
25	provide the application and information required

1	under section $351(1)(2)(A)$ of such Act, an applica-
2	tion seeking approval of a biological product for a
3	patent that could be identified pursuant to section
4	351(l)(3)(A)(i) of such Act,"; and
5	(iv) in the matter following subpara-
6	graph (C) (as added by clause (iii)), by
7	striking "or veterinary biological product"
8	and inserting ", veterinary biological prod-
9	uct, or biological product";
10	(B) in paragraph (4)—
11	(i) in subparagraph (B), by—
12	(I) striking "or veterinary bio-
13	logical product" and inserting ", vet-
14	erinary biological product, or biologi-
15	cal product"; and
16	(II) striking "and" at the end;
17	(ii) in subparagraph (C), by—
18	(I) striking "or veterinary bio-
19	logical product" and inserting ", vet-
20	erinary biological product, or biologi-
21	cal product"; and
22	(II) striking the period and in-
23	serting ", and";
24	(iii) by inserting after subparagraph
25	(C) the following:

1	"(D) the court shall order a permanent injunc-
2	tion prohibiting any infringement of the patent by
3	the biological product involved in the infringement
4	until a date which is not earlier than the date of the
5	expiration of the patent that has been infringed
6	under paragraph (2)(C), provided the patent is the
7	subject of a final court decision, as defined in sec-
8	tion 351(k)(6) of the Public Health Service Act, in
9	an action for infringement of the patent under sec-
10	tion 351(l)(6) of such Act, and the biological prod-
11	uct has not yet been approved because of section
12	351(k)(7) of such Act."; and
13	(iv) in the matter following subpara-
14	graph (D) (as added by clause (iii)), by
15	striking "and (C)" and inserting "(C), and
16	(D)"; and
17	(C) by adding at the end the following:
18	"(6)(A) Subparagraph (B) applies, in lieu of para-
19	graph (4), in the case of a patent—
20	"(i) that is identified, as applicable, in the list
21	of patents described in section 351(l)(4) of the Pub-
22	lic Health Service Act or the lists of patents de-
23	scribed in section 351(l)(5)(B) of such Act with re-
24	spect to a biological product; and

1	"(ii) for which an action for infringement of the
2	patent with respect to the biological product—
3	"(I) was brought after the expiration of
4	the 30-day period described in subparagraph
5	(A) or (B), as applicable, of section 351(l)(6) of
6	such Act; or
7	"(II) was brought before the expiration of
8	the 30-day period described in subclause (I)
9	but which was dismissed without prejudice or
10	was not prosecuted to judgment in good faith
11	"(B) In an action for infringement of a patent de-
12	scribed in subparagraph (A), the sole and exclusive remedy
13	that may be granted by a court, upon a finding that the
14	making, using, offering to sell, selling, or importation into
15	the United States of the biological product that is the sub-
16	ject of the action infringed the patent, shall be a reason-
17	able royalty.
18	"(C) The owner of a patent that should have been
19	included in the list described in section 351(l)(3)(A) of
20	the Public Health Service Act, including as provided under
21	section 351(l)(7) of such Act for a biological product, but
22	was not timely included in such list, may not bring an
23	action under this section for infringement of the patent
24	with respect to the biological product.".

24 tion.".

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1	(2) Conforming amendment under title
2	28.—Section 2201(b) of title 28, United States
3	Code, is amended by inserting before the period the
4	following: ", or section 351 of the Public Health
5	Service Act".
6	(d) Conforming Amendments Under the Fed-
7	ERAL FOOD, DRUG, AND COSMETIC ACT.—
8	(1) Content and review of applica-
9	TIONS.—Section 505(b)(5)(B) of the Federal Food,
10	Drug, and Cosmetic Act (21 U.S.C. 355(b)(5)(B)) is
11	amended by inserting before the period at the end
12	of the first sentence the following: "or, with respect
13	to an applicant for approval of a biological product
14	under section 351(k) of the Public Health Service
15	Act, any necessary clinical study or studies".
16	(2) New active ingredient.—Section 505B
17	of the Federal Food, Drug, and Cosmetic Act (21
18	U.S.C. 355c) is amended by adding at the end the
19	following:
20	"(i) New Active Ingredient.—A biological prod-
21	uct that is interchangeable with a reference product under
22	section 351 of the Public Health Service Act shall not be
23	considered to have a new active ingredient under this sec-
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1	(e) Products Previously Approved Under Sec-
2	TION 505.—
3	(1) Requirement to follow section 351.—
4	Except as provided in paragraph (2), an application
5	for a biological product shall be submitted under
6	section 351 of the Public Health Service Act (42
7	U.S.C. 262) (as amended by this Act).
8	(2) Exception.—An application for a biologi-
9	cal product may be submitted under section 505 of
10	the Federal Food, Drug, and Cosmetic Act (21
11	U.S.C. 355) if—
12	(A) such biological product is in a product
13	class for which a biological product in such
14	product class is the subject of an application
15	approved under such section 505 not later than
16	the date of enactment of this Act; and
17	(B) such application—
18	(i) has been submitted to the Sec-
19	retary of Health and Human Services (re-
20	ferred to in this Act as the "Secretary")
21	before the date of enactment of this Act;
22	or
23	(ii) is submitted to the Secretary not
24	later than the date that is 10 years after
25	the date of enactment of this Act.

- 1 (3) LIMITATION.—Notwithstanding paragraph 2 (2), an application for a biological product may not 3 be submitted under section 505 of the Federal Food, 4 Drug, and Cosmetic Act (21 U.S.C. 355) if there is 5 another biological product approved under sub-6 section (a) of section 351 of the Public Health Serv-7 ice Act that could be a reference product with re-8 spect to such application (within the meaning of 9 such section 351) if such application were submitted 10 under subsection (k) of such section 351. 11 (4)DEEMED APPROVED UNDER SECTION 12 351.—An approved application for a biological prod-13 uct under section 505 of the Federal Food, Drug, 14 and Cosmetic Act (21 U.S.C. 355) shall be deemed 15 to be a license for the biological product under such 16 section 351 on the date that is 10 years after the 17 date of enactment of this Act. 18 (5) Definitions.—For purposes of this sub-19 section, the term "biological product" has the mean-20 ing given such term under section 351 of the Public 21 Health Service Act (42 U.S.C. 262) (as amended by 22 this Act).
- 23 (f) Follow-on Biologics User Fees.—
- 24 (1) Development of user fees for bio-25 Similar biological products.—

1	(A) IN GENERAL.—Beginning not later
2	than October 1, 2010, the Secretary shall de-
3	velop recommendations to present to Congress
4	with respect to the goals, and plans for meeting
5	the goals, for the process for the review of bio-
6	similar biological product applications sub-
7	mitted under section 351(k) of the Public
8	Health Service Act (as added by this Act) for
9	the first 5 fiscal years after fiscal year 2012. In
10	developing such recommendations, the Sec-
11	retary shall consult with—
12	(i) the Committee on Health, Edu-
13	cation, Labor, and Pensions of the Senate;
14	(ii) the Committee on Energy and
15	Commerce of the House of Representa-
16	tives;
17	(iii) scientific and academic experts;
18	(iv) health care professionals;
19	(v) representatives of patient and con-
20	sumer advocacy groups; and
21	(vi) the regulated industry.
22	(B) Public review of recommenda-
23	TIONS.—After negotiations with the regulated
24	industry, the Secretary shall—

1	(i) present the recommendations de-
2	veloped under subparagraph (A) to the
3	Congressional committees specified in such
4	subparagraph;
5	(ii) publish such recommendations in
6	the Federal Register;
7	(iii) provide for a period of 30 days
8	for the public to provide written comments
9	on such recommendations;
10	(iv) hold a meeting at which the pub-
11	lic may present its views on such rec-
12	ommendations; and
13	(v) after consideration of such public
14	views and comments, revise such rec-
15	ommendations as necessary.
16	(C) Transmittal of Recommenda-
17	TIONS.—Not later than January 15, 2012, the
18	Secretary shall transmit to Congress the revised
19	recommendations under subparagraph (B), a
20	summary of the views and comments received
21	under such subparagraph, and any changes
22	made to the recommendations in response to
23	such views and comments.
24	(2) Establishment of user fee pro-
25	GRAM.—It is the sense of the Senate that, based on

- the recommendations transmitted to Congress by the
 Secretary pursuant to paragraph (1)(C), Congress
 should authorize a program, effective on October 1,
 2012, for the collection of user fees relating to the
 submission of biosimilar biological product applications under section 351(k) of the Public Health
 Service Act (as added by this Act).
 - (3) Transitional provisions for user fees for biosimilar biological products.—
 - (A) APPLICATION OF THE PRESCRIPTION DRUG USER FEE PROVISIONS.—Section 735(1)(C) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 379g(1)(C)) is amended by striking "section 351" and inserting "subsection (a) or (k) of section 351".
 - (B) EVALUATION OF COSTS OF REVIEWING BIOSIMILAR BIOLOGICAL PRODUCT APPLICATIONS.—During the period beginning on the date of enactment of this Act and ending on October 1, 2010, the Secretary shall collect and evaluate data regarding the costs of reviewing applications for biological products submitted under section 351(k) of the Public Health Service Act (as added by this Act) during such period.

1	(C) AUDIT.—
2	(i) IN GENERAL.—On the date that is
3	2 years after first receiving a user fee ap-
4	plicable to an application for a biological
5	product under section 351(k) of the Public
6	Health Service Act (as added by this Act).
7	and on a biennial basis thereafter until Oc-
8	tober 1, 2013, the Secretary shall perform
9	an audit of the costs of reviewing such ap-
10	plications under such section 351(k). Such
11	an audit shall compare—
12	(I) the costs of reviewing such
13	applications under such section
14	351(k) to the amount of the user fee
15	applicable to such applications; and
16	(II)(aa) such ratio determined
17	under subclause (I); to
18	(bb) the ratio of the costs of re-
19	viewing applications for biological
20	products under section 351(a) of such
21	Act (as amended by this Act) to the
22	amount of the user fee applicable to
23	such applications under such section
24	351(a).

1	(ii) Alteration of user fee.—If
2	the audit performed under clause (i) indi-
3	cates that the ratios compared under sub-
4	clause (II) of such clause differ by more
5	than 5 percent, then the Secretary shall
6	alter the user fee applicable to applications
7	submitted under such section 351(k) to
8	more appropriately account for the costs of
9	reviewing such applications.
10	(iii) Accounting standards.—The
11	Secretary shall perform an audit under
12	clause (i) in conformance with the account-
13	ing principles, standards, and requirements
14	prescribed by the Comptroller General of
15	the United States under section 3511 of
16	title 31, United State Code, to ensure the
17	validity of any potential variability.
18	(4) Authorization of appropriations.—
19	There is authorized to be appropriated to carry out
20	this subsection such sums as may be necessary for
21	each of fiscal years 2008 through 2012.
22	(g) Allocation of Savings; Special Reserve
23	Fund.—
24	(1) Determination of savings.—The Sec-
25	retary of the Treasury, in consultation with the Sec-

1	retary, shall for each fiscal year determine the
2	amount of the savings to the Federal Government as
3	a result of the enactment of this Act and shall trans-
4	fer such amount to the Fund established under
5	paragraph (2) pursuant to a relevant appropriations
6	Act.

(2) Special reserve fund.—

- (A) IN GENERAL.—There is established in the Treasury of the United States a fund to be designated as the "Biological Product Savings Fund" to be made available to the Secretary without fiscal year limitation.
- (B) USE OF FUND.—The amounts made available to the Secretary through the Fund under subparagraph (A) shall be expended on activities authorized under the Public Health Service Act.
- (3) AUTHORIZATION OF APPROPRIATIONS.—
 There is authorized to be appropriated for each fiscal year to the Fund established under paragraph (2), the amount of the savings determined for such fiscal year under paragraph (1).