[116H2387]

(Origina	al Signature	of Member)

117TH CONGRESS 1ST SESSION

H. R. 2843

To amend subsection (q) of section 505 of the Federal Food, Drug, and Cosmetic Act to clarify the process for denying certain petitions whose primary purpose is to delay the approval of an application submitted under subsection (b)(2) or (j) of such section 505, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

Mr. LEVII	N of Michigan in	troduced the	e following	bill; whic	h was	referred	to
	the Committee	on					

A BILL

To amend subsection (q) of section 505 of the Federal Food, Drug, and Cosmetic Act to clarify the process for denying certain petitions whose primary purpose is to delay the approval of an application submitted under subsection (b)(2) or (j) of such section 505, 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,

1 SECTION 1. SHORT TITLE.

2	This Act may be cited as the "Stop The Overuse of
3	Petitions and Get Affordable Medicines to Enter Soon Act
4	of 2021" or the "STOP GAMES Act of 2021".
5	SEC. 2. DENIAL OF PETITIONS WHOSE PRIMARY PURPOSE
6	IS TO DELAY APPROVAL OF CERTAIN APPLI-
7	CATIONS.
8	(a) In General.—Subparagraph (E) of section
9	505(q)(1) of the Federal Food, Drug, and Cosmetic Act
10	$(21~\mathrm{U.S.C.}~355(q)(1))$ is amended to read as follows:
11	"(E) Denial based on intent to
12	DELAY.—
13	"(i) In General.—If the Secretary
14	determines that a petition or a supplement
15	to the petition was submitted with the pri-
16	mary purpose of delaying the approval of
17	an application or the petition does not on
18	its face raise valid scientific or regulatory
19	issues, the Secretary may deny the petition
20	at any point based on such determination.
21	"(ii) Factors.—The Secretary may
22	issue guidance to describe the factors that
23	will be used to determine under this sub-
24	paragraph whether a petition is submitted
25	with the primary purpose of delaying the

1	approval of an application. Such factors
2	shall include the following:
3	"(I) Submission of a petition
4	where it appears, based on the date
5	that relevant information relied upon
6	in the petition became known to the
7	petitioner (or reasonably should have
8	been known to the petitioner), that
9	the petitioner has taken an unreason-
10	able length of time to submit the peti-
11	tion.
12	"(II) Submission of multiple or
13	serial petitions raising issues that rea-
14	sonably could have been known to the
15	petitioner at the time of submission of
16	the earlier petition or petitions.
17	"(III) Submission of a petition
18	close in time to a known, first date
19	upon which an application under sub-
20	section (b)(2) or (j) of this section or
21	under section 351(k) of the Public
22	Health Service Act could be approved
23	(such as submission close in time to
24	the expiration of a blocking patent or
25	exclusivity).

1	"(IV) Submission of a petition
2	without any data or information in
3	support of the scientific positions set
4	forth in the petition.
5	"(V) Submission of a petition
6	raising the same or substantially simi-
7	lar issues as a prior petition to which
8	the Food and Drug Administration
9	has already substantively responded,
10	particularly where the subsequent sub-
11	mission closely follows in time the ear-
12	lier response.
13	"(VI) Submission of a petition
14	concerning standards for approval of
15	a drug product for which—
16	"(aa) the Food and Drug
17	Administration has provided an
18	opportunity for public input
19	(such as when the Food and
20	Drug Administration has issued
21	draft or final product-specific
22	guidance applicable to the drug
23	product); and

1	"(bb) the petitioner has not
2	provided comment other than
3	through the petition.
4	"(VII) Submission of a petition
5	requesting that other applicants must
6	meet standards for testing, data, or
7	labeling for their products that are
8	more onerous or rigorous than the
9	standards applicable to the applicable
10	listed drug or the petitioner's version
11	of the same product.
12	"(VIII) Other relevant consider-
13	ations, including the history of the pe-
14	titioner with the Food and Drug Ad-
15	ministration (such as whether the pe-
16	titioner has a history of submitting
17	petitions which the Food and Drug
18	Administration has determined were
19	submitted with the primary purpose of
20	delay).
21	"(iii) Referral to ftc.—If the Sec-
22	retary determines that a petition has been
23	submitted with the primary purpose of de-
24	laying the approval of an application, as
25	described in clause (i), the Secretary shall

1	refer the matter to the Federal Trade
2	Commission.".
3	(b) Deadline for Submission of Petitions.—
4	(1) Deadline.—Clause (i) of section
5	505(q)(1)(A) of the Federal Food, Drug, and Cos-
6	metic Act (21 U.S.C. 355(q)(1)(A)) is amended to
7	read as follows:
8	"(i) the request is in writing, is a pe-
9	tition submitted to the Secretary pursuant
10	to section 10.30, 10.31, or 10.35 of title
11	21, Code of Federal Regulations (or any
12	successor regulations), and is submitted
13	not later than 60 days after the informa-
14	tion upon which the petition is based first
15	became known to the party on whose be-
16	half the petition is submitted; and".
17	(2) Certification.—Section 505(q)(1)(H) of
18	the Federal Food, Drug, and Cosmetic Act (21
19	U.S.C. 355(q)(1)) is amended by striking "I further
20	certify that the information upon which I have based
21	the action requested herein first became known to
22	the party on whose behalf this petition is submitted
23	on or about the following date:" and in-
24	serting "I further certify that the information upon
25	which I have based the action requested herein first

1	became known to the party on whose behalf this pe-
2	tition is submitted on or about, which
3	date was not more than 60 days before the date of
4	submitting this petition.".
5	(c) Reporting to Congress.—Section 505(q)(3) of
6	the Federal Food, Drug, and Cosmetic Act (21 U.S.C.
7	355(q)(3)) is amended—
8	(1) in the matter before subparagraph (A), by
9	striking "specifies";
10	(2) in subparagraphs (A), (B), (C), and (D), by
11	striking "the number" and inserting "specifies the
12	number";
13	(3) in subparagraph (C), by striking "and" at
14	the end;
15	(4) in subparagraph (D), by striking the period
16	at the end and inserting "; and; and
17	(5) by adding at the end the following:
18	"(E)(i) lists each petition submitted during
19	such period and, for each, identifies the peti-
20	tioner;
21	"(ii) quantifies the time and resources ex-
22	pended on each such petition;
23	"(iii) states the timing of the petition rel-
24	ative to the expiration date of the patents speci-
25	fied in the pending application in the certifi-

1	cation under subsection $(b)(2)(A)$ or
2	(j)(2)(A)(vii), as applicable;
3	"(iv) quantifies the delay, if any, caused by
4	any such petition on the approval of any appli-
5	cation submitted under subsection (b)(2) or (j),
6	including a description of how any such delay is
7	calculated and an estimate of when any delayed
8	approval would have been granted absent the
9	petition; and
10	"(v) in cases in which a pending applica-
11	tion and a petition with respect to such pending
12	application are disposed of on the same or near-
13	ly the same date, states when the Food and
14	Drug Administration would have disposed of
15	the pending application absent the petition.".