H. R. 6972

To amend the Federal Food, Drug, and Cosmetic Act to establish additional authorities of the Food and Drug Administration regarding the conduct of pediatric investigations of molecularly targeted drugs to treat cancer, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

Mr. BUTTERFIELD introduced the following bill; which was referred to the Committee on

A BILL

To amend the Federal Food, Drug, and Cosmetic Act to establish additional authorities of the Food and Drug Administration regarding the conduct of pediatric investigations of molecularly targeted drugs to treat cancer, and for other purposes.

Be it enacted by the Senate and House of Representa-
tives of the United States of America in Congress assembled,

SECTION 1. SHORT TITLE.

This Act may be cited as the “Give Kids a Chance
Act of 2022”.

March 8, 2022 (2:19 p.m.)
SEC. 2. RESEARCH INTO PEDIATRIC USES OF DRUGS; ADDITIONAL AUTHORITIES OF FOOD AND DRUG ADMINISTRATION REGARDING MOLECULARLY TARGETED CANCER DRUGS.

(a) **IN GENERAL.**—

(1) **AUTHORITY REGARDING INVESTIGATION OF NOVEL COMBINATION DRUGS.**—Section 505B(a)(1)(B) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355c(a)(1)(B)) is amended, in the matter preceding clause (i), by inserting after “Public Health Service Act,” the following: “or an application under such section 505 or such section 351 for a drug or biological product that contains a novel combination of two or more active ingredients (subject to paragraph (3)(B)(iii))”

(2) **ADDITIONAL ACTIVE INGREDIENT FOR APPLICATION DRUG; LIMITATION REGARDING NOVEL-COMBINATION APPLICATION DRUG.**—Section 505B(a)(3) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355c(a)(3)) is amended—

(A) by redesignating subparagraphs (B) and (C) as subparagraphs (C) and (D), respectively; and

(B) by striking subparagraph (A) and inserting the following:
“(A) IN GENERAL.—For purposes of paragraph (1)(B), the investigation described in this paragraph is (as determined by the Secretary) a molecularly targeted pediatric cancer investigation of—

“(i) the drug or biological product for which the application referred to in such paragraph is submitted; or

“(ii) the active ingredient or ingredients of such drug or biological product in combination with—

“(I) an active ingredient of a drug for which an approved application under section 505(j) is in effect or an active ingredient of a biological product for which an approved application under section 351(k) of the Public Health Service Act is in effect, which drug or biological product is determined by the Secretary to be the standard of care for treating a pediatric cancer;

“(II) an active ingredient of a drug for which an approved application under section 505(b) is in effect
to treat an adult cancer, or an active
ingredient of a biological product for
which an approved application under
section 351(a) of the Public Health
Service Act is in effect to treat an
adult cancer, which approved applica-
tion is held by the same person sub-
mitting the application referred to in
paragraph (1)(B); or

“(III) an active ingredient of a
drug or biological product for which
there is in effect an exemption for in-
vestigational use under section 505(i),
which drug or biological product is
under such exemption being studied
jointly by the person submitting the
application referred to in paragraph
(1)(B) and by another person pursu-
ant to an agreement between such
persons.

“(B) ADDITIONAL REQUIREMENTS.—

“(i) DESIGN OF INVESTIGATION.—A
molecularly targeted pediatric cancer inves-
tigation referred to in subparagraph (A)
shall be designed to yield clinically mean-
ingful pediatric study data, gathered using appropriate formulations for each age group for which the study is required, re-
garding dosing, safety, and preliminary ef-
ficacy.

“(ii) PURPOSE OF INVESTIGATION.—
The purpose of a molecularly targeted pe-
diatric cancer investigation referred to in subparagraph (A) shall be—

“(I) in the case of such an invest-
tigation conducted with respect to a
drug or biological product referred to in clause (i) of such subparagraph, to inform potential pediatric labeling of the drug or biological product for
which the application referred to in paragraph (1)(B) is submitted; and

“(II) in the case of such an in-
vestigation conducted with respect to a combination of active ingredients
described to in clause (ii) of such sub-
paragraph, to assist in determining
the relevance of its molecular target
to the growth or progression of a pe-
diatric cancer.
“(iii) LIMITATION REGARDING INVESTIGATION OF NOVEL COMBINATION.—For purposes of paragraph (1)(B), a novel combination is a combination of two or more active ingredients for which an application under section 505 of this Act or section 351 of the Public Health Service Act for such combination has not previously been approved. A pediatric investigation under this paragraph of such novel combination is required only if each of the active ingredients in the combination has been approved under such section 505 or such section 351 to treat an adult cancer.

“(iv) PRECLINICAL DATA.—The Secretary may require that reports on an investigation required pursuant to paragraph (1)(B) shall include the results of all preclinical studies on which the decision to conduct such investigation was based.

“(v) RULE OF CONSTRUCTION REGARDING INACTIVE INGREDIENTS.—With respect to a combination of active ingredients referred to in subparagraph (A)(ii), such subparagraph may not be construed
as addressing the use of inactive ingredients with such combination.”.

(3) CLARIFYING APPLICABILITY OF CERTAIN PROVISIONS.—Section 505B(a)(3) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355e(a)(3)), as amended by paragraph (2), is further amended by adding at the end the following:

“(E) INTERNAL COMMITTEE REVIEW; LABELING CHANGES; DISSEMINATION OF INFORMATION; ADVERSE EVENTS; SCOPE OF AUTHORITY.—Subsections (f) through (j) shall apply with respect to investigations described in this paragraph to the same extent and in the same manner as such subsections apply with respect to the assessments required under paragraph (1)(A), except that subsection (g) does not apply with respect to an investigation referred to in subparagraph (A)(ii) of this paragraph.”.

(4) CONFORMING AMENDMENTS.—Section 505B(a) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355e(a)) is amended—

(A) in paragraph (3)(C), as redesignated by paragraph (2)(A) of this subsection, by striking “investigations described in this para-
graph” and inserting “investigations referred to in subparagraph (A)(i)”;

(B) in paragraph (3)(D), as redesignated by paragraph (2)(A) of this subsection, by striking “the assessments under paragraph (2)(B)” and inserting “the assessments required under paragraph (1)(A)”;

and

(C) in paragraph (5)(D), by inserting before the period at the end the following: “, except this subparagraph is not applicable to an investigation referred to in paragraph (3)(A)(ii)”.

(b) Authority Regarding Preclinical Studies.—Section 505B(a)(1) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355e(a)(1)), as amended by subsection (a)(1), is further amended by adding at the end the following:

“(C) Preclinical Studies Generally.—

“(i) In general.—With respect to an application for an exemption for investigational use under section 505(i) for a drug or biological product that is intended for the treatment of an adult cancer, the Secretary may require, as a condition of per-
mitting the exemption to go into effect, that the sponsor involved enter into an agreement with the Secretary to conduct not more than two preclinical studies of the drug or biological product in order to assist in determining the relevance of its molecular target to the growth or progression of a pediatric cancer.

“(ii) **Timeframe for Preclinical Studies.**—With respect to the drug or biological product involved, an agreement under clause (i) for a preclinical study shall specify the date by which an initial plan for the study will be submitted to the Secretary except that the Secretary may not require the submission of such plan any earlier than one year after the exemption referred to in clause (i) goes into effect. The results of the preclinical study shall be submitted to the Secretary in accordance with a timeframe to which the Secretary and the sponsor involved have agreed. Such timeframe shall provide for deferrals equivalent to deferrals under paragraph (4).”
(c) APPLICABILITY.—The amendments made by this section apply with respect to any application under section 505(i) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355(i)), any application under section 505 of such Act (21 U.S.C. 355), and any application under section 351(a) of the Public Health Service Act (42 U.S.C. 262), that is submitted on or after the expiration of the 3-year period beginning on the date of the enactment of this Act.