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(Original Signature of Member)

115TH CONGRESS
2D SESSION

H. R. _____

To amend title XVIII of the Social Security Act to require the Secretary of Health and Human Services to negotiate prices of prescription drugs furnished under part D of the Medicare program.

IN THE HOUSE OF REPRESENTATIVES

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

A BILL

To amend title XVIII of the Social Security Act to require the Secretary of Health and Human Services to negotiate prices of prescription drugs furnished under part D of the Medicare program.

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 “Medicare Negotiation
5 and Competitive Licensing Act of 2018”.

1 **SEC. 2. REQUIRING THE SECRETARY OF HEALTH AND**
2 **HUMAN SERVICES TO NEGOTIATE PRICES OF**
3 **PRESCRIPTION DRUGS FURNISHED UNDER**
4 **PART D OF THE MEDICARE PROGRAM.**

5 Section 1860D–11 of the Social Security Act (42
6 U.S.C. 1395w–111) is amended by striking subsection (i)
7 and inserting the following new subsection:

8 “(i) NEGOTIATION OF LOWER DRUG PRICES.—

9 “(1) IN GENERAL.—Notwithstanding any other
10 provision of law, the Secretary shall, for plan years
11 beginning on or after the date of the enactment of
12 this subsection, negotiate with pharmaceutical man-
13 ufacturers the prices (including discounts, rebates,
14 and other price concessions) that may be charged to
15 PDP sponsors and MA organizations during a nego-
16 tiated price period (as specified by the Secretary) for
17 covered part D drugs for part D eligible individuals
18 who are enrolled under a prescription drug plan or
19 under an MA-PD plan. In negotiating such prices
20 under this section, the Secretary shall take into ac-
21 count the following factors:

22 “(A) The comparative clinical effectiveness
23 and cost effectiveness, when available from an
24 impartial source, of such drug.

25 “(B) The budgetary impact of providing
26 coverage of such drug.

1 “(C) The number of similarly effective
2 drugs or alternative treatment regimens for
3 each approved use of such drug.

4 “(D) The associated financial burden on
5 patients that utilize such drug.

6 “(E) The associated unmet patient need
7 for such drug.

8 “(F) The total revenues from global sales
9 obtained by the manufacturer for such drug.

10 “(2) FINALIZATION OF NEGOTIATED PRICE.—

11 The negotiated price of each covered part D drug for
12 a negotiated price period shall be finalized not later
13 than 30 days before a PDP sponsor is required to
14 submit information described in subsection (b)(2)
15 for the first plan year in such negotiated price pe-
16 riod.

17 “(3) COMPETITIVE LICENSING AUTHORITY.—

18 “(A) IN GENERAL.—Notwithstanding any
19 exclusivity under clause (iii) or (iv) of section
20 505(j)(5)(F) of the Federal Food, Drug, and
21 Cosmetic Act, clause (iii) or (iv) of section
22 505(c)(3)(E) of such Act, section 351(k)(7)(A)
23 of the Public Health Service Act, or section
24 527(a) of the Federal Food, Drug, and Cos-
25 metic Act, or by an extension of such exclusivity

1 under section 505A of such Act or section 505E
2 of such Act, and any other provision of law that
3 provides for market exclusivity (or extension of
4 market exclusivity) with respect to a drug, in
5 the case that the Secretary is unable to success-
6 fully negotiate an appropriate price for a cov-
7 ered part D drug for a negotiated price period,
8 the Secretary shall authorize the use of any
9 patent, clinical trial data, or other exclusivity
10 granted by the Federal government with respect
11 to such drug as the Secretary determines ap-
12 propriate for purposes of manufacturing such
13 drug for sale under a prescription drug plan or
14 MA-PD plan. Any entity making use of a com-
15 petitive license to use patent, clinical trial data,
16 or other exclusivity under this section shall pro-
17 vide to the manufacturer holding such exclu-
18 sivity reasonable compensation, as determined
19 by the Secretary based on the following factors:

20 “(i) The risk-adjusted value of any
21 Federal government subsidies and invest-
22 ments in research and development used to
23 support the development of such drug.

24 “(ii) The risk-adjusted value of any
25 investment made by such manufacturer in

1 the research and development of such
2 drug.

3 “(iii) The impact of the price, includ-
4 ing license compensation payments, on
5 meeting the medical need of all patients.

6 “(iv) The relationship between the
7 price of such drug, including compensation
8 payments, and the health benefits of such
9 drug.

10 “(v) Other relevant factors determined
11 appropriate by the Secretary to provide
12 reasonable compensation.

13 “(B) REASONABLE COMPENSATION.—The
14 manufacturer described in subparagraph (A)
15 may seek recovery against the United States in
16 the United States Court of Federal Claims.

17 “(C) INTERIM PERIOD.—

18 “(i) IN GENERAL.—Until a drug de-
19 scribed in subparagraph (A) is approved
20 under section 505(j) of the Federal Food,
21 Drug, and Cosmetic Act or section 351(k)
22 of the Public Health Service Act and is
23 provided under license issued by the Sec-
24 retary under such subparagraph, PDP
25 plans and MA-PD plans shall not pay

1 more for such drug than the average of the
2 prices available, during the most recent 12-
3 month period for which data is available
4 prior to the beginning of such negotiated
5 price period, from the manufacturer to any
6 wholesaler, retailer, provider, health main-
7 tenance organization, nonprofit entity, or
8 governmental entity in the ten OECD (Or-
9 ganization for Economic Cooperation and
10 Development) countries that have the larg-
11 est gross domestic product with a per cap-
12 ita income that is not less than half the
13 per capita income of the United States, as
14 reported by the manufacturer to the Sec-
15 retary.

16 “(ii) FEDERAL PROGRAM LICENS-
17 ING.—If such drug is not made available
18 at the price determined, the Secretary shall
19 authorize such entities to use any patent,
20 clinical trial data, or other exclusivity
21 granted by the Federal government with
22 respect to such drug as the Secretary de-
23 termines appropriate for purposes of man-
24 ufacturing such drug for sale under any
25 Federal program, including those provided

1 by Medicare, Medicaid, Veterans Affairs,
2 the Department of Defense, and the Coast
3 Guard.

4 “(4) FDA EXPEDITED REVIEW OF LICENSED
5 DRUG APPLICATIONS.—The Secretary shall prioritize
6 review of applications under section 505(j) of the
7 Federal Food, Drug, and Cosmetic Act for drugs li-
8 censed under paragraph (3)(A).

9 “(5) PROHIBITION OF ANTICOMPETITIVE BE-
10 HAVIOR.—No drug manufacturer may engage in
11 anticompetitive behavior with another manufacturer
12 that may interfere with the issuance and implemen-
13 tation of a competitive license or run contrary to
14 public policy.

15 “(6) CLARIFICATION.—Nothing in this sub-
16 section shall be construed as preventing the sponsor
17 of a prescription drug plan or an organization offer-
18 ing an MA-PD plan from obtaining a discount or re-
19 duction of the price for a covered part D drug below
20 the price negotiated by the Secretary.”.