

115TH CONGRESS  
1ST SESSION

# H. R. 3528

To amend title XVIII of the Social Security Act to require e-prescribing for coverage under part D of the Medicare program of prescription drugs that are controlled substances.

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## IN THE HOUSE OF REPRESENTATIVES

JULY 28, 2017

Ms. CLARK of Massachusetts (for herself and Mr. MULLIN) introduced the following bill; which was referred to the Committee on Energy and Commerce, and in addition to the Committee on Ways and Means, for a period to be subsequently determined by the Speaker, in each case for consideration of such provisions as fall within the jurisdiction of the committee concerned

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## A BILL

To amend title XVIII of the Social Security Act to require e-prescribing for coverage under part D of the Medicare program of prescription drugs that are controlled substances.

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 “Every Prescription  
5 Conveyed Securely Act”.

1 **SEC. 2. REQUIRING E-PRESCRIBING FOR COVERAGE OF**  
2 **COVERED PART D CONTROLLED SUB-**  
3 **STANCES.**

4 (a) IN GENERAL.—Section 1860D–4(e) of the Social  
5 Security Act (42 U.S.C. 1395w–104(e)) is amended by  
6 adding at the end the following:

7 “(7) REQUIREMENT OF E-PRESCRIBING FOR  
8 CONTROLLED SUBSTANCES.—

9 “(A) IN GENERAL.—Subject to subpara-  
10 graph (B), a prescription for a covered part D  
11 drug under a prescription drug plan (or under  
12 an MA–PD plan) for a schedule II, III, IV, or  
13 V controlled substance shall be transmitted by  
14 a health care practitioner electronically in ac-  
15 cordance with an electronic prescription drug  
16 program that meets the requirements of para-  
17 graph (2).

18 “(B) EXCEPTION FOR CERTAIN CIR-  
19 CUMSTANCES.—The Secretary shall, pursuant  
20 to rulemaking, specify circumstances with re-  
21 spect to which the Secretary may waive the re-  
22 quirement under subparagraph (A), with re-  
23 spect to a covered part D drug, including in the  
24 case of—

1           “(i) a prescription issued when the  
2           prescriber and dispenser are the same enti-  
3           ty;

4           “(ii) a prescription issued that cannot  
5           be transmitted electronically due to the  
6           constraints of the most recently imple-  
7           mented version of the National Council for  
8           Prescription Drug Programs SCRIPT  
9           Standard;

10          “(iii) a prescription issued by a practi-  
11          tioner who has received a waiver or a re-  
12          newal thereof for a specified period deter-  
13          mined by the Secretary, not to exceed one  
14          year, from the requirement to use elec-  
15          tronic prescribing, pursuant to a process  
16          established by regulation by the Secretary,  
17          due to demonstrated economic hardship,  
18          technological limitations that are not rea-  
19          sonably within the control of the practi-  
20          tioner, or other exceptional circumstance  
21          demonstrated by the practitioner;

22          “(iv) a prescription issued by a practi-  
23          tioner under circumstances in which, not-  
24          withstanding the practitioner’s ability to  
25          make an electronic prescription as required

1 by this subsection, such practitioner rea-  
2 sonably determines that it would be im-  
3 practical for the individual involved to ob-  
4 tain substances prescribed by electronic  
5 prescription in a timely manner, and such  
6 delay would adversely impact the individ-  
7 ual’s medical condition involved;

8 “(v) a prescription issued by a practi-  
9 tioner allowing for the dispensing of a non-  
10 patient specific prescription pursuant to a  
11 standing order, approved protocol for drug  
12 therapy, collaborative drug management,  
13 or comprehensive medication management,  
14 in response to a public health emergency,  
15 or other circumstances where the practi-  
16 tioner may issue a non-patient specific pre-  
17 scription;

18 “(vi) a prescription issued by a practi-  
19 tioner prescribing a drug under a research  
20 protocol; and

21 “(vii) a prescription issued by a prac-  
22 titioner for a drug for which the Food and  
23 Drug Administration requires the prescrip-  
24 tion to contain certain elements that are  
25 not able to be accomplished with electronic

1           prescribing such as, a drug with risk eval-  
2           uation and mitigation strategies that in-  
3           clude elements to assure safe use.

4           “(C) DISPENSING.—Nothing in this para-  
5           graph shall be construed as requiring a sponsor  
6           of a prescription drug plan under this part, MA  
7           organization offering an MA–PD plan under  
8           part C, or a pharmacist to verify that a practi-  
9           tioner, with respect to a prescription for a cov-  
10          ered part D drug, has a waiver (or is otherwise  
11          exempt) under subparagraph (B) from the re-  
12          quirement under subparagraph (A). Nothing in  
13          this paragraph shall be construed as affecting  
14          the ability of the plan to cover or the phar-  
15          macists’ ability to continue to dispense covered  
16          part D drugs from otherwise valid written, oral  
17          or fax prescriptions that are consistent with  
18          laws and regulations.

19          “(D) ENFORCEMENT.—The Secretary  
20          shall, pursuant to rulemaking, have authority to  
21          enforce and specify appropriate penalties for  
22          non-compliance with the requirement under  
23          subparagraph (A).”.

1           (b) EFFECTIVE DATE.—The amendment made by  
2 subsection (a) shall apply to coverage of drugs prescribed  
3 on or after January 1, 2020.

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