

117TH CONGRESS
2D SESSION

S. 4351

To amend the Federal Food, Drug, and Cosmetic Act to clarify the conditions under which the Secretary of Health and Human Services can approve generic drug applications with labeling temporarily different than the brand name drug, and for other purposes.

IN THE SENATE OF THE UNITED STATES

JUNE 6, 2022

Mr. ROMNEY (for himself, Ms. HASSAN, Ms. ROSEN, and Mr. HICKENLOOPER) introduced the following bill; which was read twice and referred to the Committee on Health, Education, Labor, and Pensions

A BILL

To amend the Federal Food, Drug, and Cosmetic Act to clarify the conditions under which the Secretary of Health and Human Services can approve generic drug applications with labeling temporarily different than the brand name drug, 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 “Enhanced Access to
5 Affordable Medicines Act of 2022”.

1 **SEC. 2. ENHANCING ACCESS TO AFFORDABLE MEDICINES.**

2 Section 505(j)(10)(A) of the Federal Food, Drug,
3 and Cosmetic Act (21 U.S.C. 355(j)(10)(A)) is amended
4 by striking clauses (i) through (iv) and inserting the fol-
5 lowing:

6 “(i) a revision to the labeling of the listed drug
7 has been approved by the Secretary within 90 days
8 of when the application is otherwise eligible for ap-
9 proval under this subsection;

10 “(ii) the sponsor of the application agrees to
11 submit revised labeling for the drug that is the sub-
12 ject of the application not later than 60 days after
13 approval under this subsection of the application;
14 and

15 “(iii) the labeling revision described under
16 clause (i) does not include a change to the ‘Warn-
17 ings’ section of the labeling.”.

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