115th CONGRESS 1st Session

S. 1052

AN ACT

To strengthen the use of patient-experience data within the benefit-risk framework for approval of new drugs.

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 "Better Empowerment
3	Now to Enhance Framework and Improve Treatments Act
4	of 2017" or the "BENEFIT Act of 2017".
5	SEC. 2. STRENGTHENING THE USE PATIENT-EXPERIENCE
6	DATA WITHIN BENEFIT-RISK FRAMEWORK.
7	Section 569C of the Federal Food, Drug, and Cos-
8	metic Act (21 U.S.C. 360bbb–8c) is amended—
9	(1) in subsection $(a)(1)$ —
10	(A) in subparagraph (A), by striking ";
11	and" and inserting a semicolon;
12	(B) in subparagraph (B), by striking the
13	period and inserting "; and"; and
14	(C) by adding at the end the following:
15	"(C) as part of the risk-benefit assessment
16	framework in the new drug approval process de-
17	scribed in section $505(d)$, considering relevant
18	patient-focused drug development data, such as
19	data from patient preference studies (benefit-
20	risk), patient reported outcome data, or patient
21	experience data, developed by the sponsor of an
22	application or another party."; and
23	(2) in subsection $(b)(1)$. by inserting ", includ-
24	ing a description of how such data and information

- 1 were considered in the risk benefit assessment de-
- 2 scribed in section 505(d)" before the period.

Passed the Senate August 3, 2017.

Attest:

Secretary.

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