115TH CONGRESS 2D SESSION

H.R.5473

AN ACT

- To direct the Secretary of Health and Human Services to update or issue one or more guidances addressing alternative methods for data collection on opioid sparing and inclusion of such data in product labeling, and for other purposes.
 - 1 Be it enacted by the Senate and House of Representa-
 - ${\it 2\ tives\ of\ the\ United\ States\ of\ America\ in\ Congress\ assembled},$

1 SECTION 1. SHORT TITLE.

2	This Act may be cited as the "Better Pain Manage-
3	ment Through Better Data Act of 2018".
4	SEC. 2. GUIDANCE ADDRESSING ALTERNATIVE AP-
5	PROACHES TO DATA COLLECTION AND LA-
6	BELING CLAIMS FOR OPIOID SPARING.
7	(a) In General.—For purposes of assisting spon-
8	sors in collecting and incorporating opioid-sparing data in
9	product labeling, the Secretary of Health and Human
10	Services (referred to in this section as the "Secretary")
11	shall conduct a public meeting and update or issue one
12	or more guidances in accordance with subsection (b).
13	(b) Guidance.—
14	(1) IN GENERAL.—The Secretary of Health and
15	Human Services, acting through the Commissioner
16	of Food and Drugs, shall update or issue one or
17	more guidances addressing—
18	(A) alternative methods for data collection
19	on opioid sparing;
20	(B) alternative methods for inclusion of
21	such data in product labeling; and
22	(C) investigations other than clinical trials,
23	including partially controlled studies and objec-
24	tive trials without matched controls such as his-
25	torically controlled analyses, open-label studies,

1	and meta-analyses, on opioid sparing for inclu-
2	sion in product labeling.
3	(2) Contents.—The guidances under para-
4	graph (1) shall address—
5	(A) innovative clinical trial designs for
6	ethically and efficiently collecting data on opioid
7	sparing for inclusion in product labeling;
8	(B) primary and secondary endpoints for
9	the reduction of opioid use while maintaining
10	adequate pain control;
11	(C) use of real world evidence, including
12	patient registries, and patient reported out-
13	comes to support inclusion of opioid-sparing
14	data in product labeling; and
15	(D) how sponsors may obtain feedback
16	from the Secretary relating to such issues prior
17	to—
18	(i) commencement of such data collec-
19	tion; or
20	(ii) the submission of resulting data to
21	the Secretary.
22	(3) Public meeting.—Prior to updating or
23	issuing the guidances required by paragraph (1), the
24	Secretary shall consult with stakeholders, including
25	representatives of regulated industry, academia, pa-

tients, and provider organizations, through a public meeting to be held not later than 12 months after the date of enactment of this Act.

(4) Timing.—The Secretary shall—

- (A) not later than 12 months after the date of the public meeting required by paragraph (3), update or issue the one or more draft guidances required by paragraph (1); and
- (B) not later than 12 months after the date on which the public comment period for such draft guidances closes, finalize such guidances.

(c) Definition.—In this section:

(1) The terms "opioid sparing" and "opioid-sparing" refer to the use of drugs or devices (as defined in section 201 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 321)) that reduce pain while enabling the reduction, replacement, or avoidance of oral opioids.

- 1 (2) The term "Secretary" means the Secretary
- of Health and Human Services.

Passed the House of Representatives June 12, 2018. Attest:

Clerk.

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