(Original Signature of Member)

115th CONGRESS 2D Session

To require the Secretary of Health and Human Services to issue guidance with respect to the expedited approval of certain drugs, and for other purposes.

H.R. 5806

IN THE HOUSE OF REPRESENTATIVES

M____ introduced the following bill; which was referred to the Committee on

A BILL

- To require the Secretary of Health and Human Services to issue guidance with respect to the expedited approval of certain drugs, 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 "21st Century Tools

5 for Pain and Addiction Treatment Act".

SEC. 2. CLARIFYING FDA REGULATION OF NON-ADDICTIVE PAIN AND ADDICTION THERAPIES.

3 (a) PUBLIC MEETINGS.—Not later than 1 year after
4 the date of enactment of this Act, the Secretary of Health
5 and Human Services, acting through the Commissioner of
6 Food and Drugs, shall hold not less than one public meet7 ing to address the challenges and barriers of developing
8 nonaddictive medical products intended to treat pain or
9 addiction, which may include—

10 (1) the application of novel clinical trial designs 11 (consistent with section 3021 of the 21st Century 12 Cures Act (Public Law 114–255)), use of real world 13 evidence (consistent with section 505F of the Fed-14 eral Food, Drug, and Cosmetic Act (21 U.S.C. 15 355g)), and use of patient experience data (con-16 sistent with section 569C of the Federal Food, 17 Drug, and Cosmetic Act (21 U.S.C. 360bbb–8c)) for 18 the development of non-addictive medical products 19 intended to treat pain or addiction; and

20 (2) the application of eligibility criteria under
21 sections 506 and 515B of the Federal Food, Drug,
22 and Cosmetic Act (21 U.S.C. 356, 360e-3) for non23 addictive medical products intended to treat pain or
24 addiction.

25 (b) GUIDANCE.—Not later than one year after the
26 public meetings are conducted under subsection (a) the

3

Secretary shall issue one or more final guidance docu ments, or update existing guidance documents, to help ad dress challenges to developing non-addictive medical prod ucts to treat pain or addiction. Such guidance documents
 shall include information regarding—

6 (1) how the Food and Drug Administration 7 may apply sections 506 and 515B of the Federal 8 Food, Drug, and Cosmetic Act (21 U.S.C. 356, 9 360e-3) to non-addictive medical products intended 10 to treat pain or addiction, including the cir-11 cumstances under which the Secretary—

12 (A) may apply the eligibility criteria under
13 such sections 506 and 515B to non-opioid or
14 non-addictive medical products intended to
15 treat pain or addiction;

16 (B) considers the risk of addiction of con17 trolled substances approved to treat pain when
18 establishing unmet medical need; and

19 (C) considers pain, pain control, or pain
20 management in assessing whether a disease or
21 condition is a serious or life-threatening disease
22 or condition; and

(2) the methods by which sponsors may evaluate acute and chronic pain, endpoints for non-addictive medical products intended to treat pain, the

4

manner in which endpoints and evaluations of efficacy will be applied across and within review divisions, taking into consideration the etiology of the
underlying disease, and the manner in which sponsors may use surrogate endpoints, intermediate
endpoints, and real world evidence.

7 (c) MEDICAL PRODUCT DEFINED.—In this section, the term "medical product" means a drug (as defined in 8 9 section 201(g)(1) of the Federal Food, Drug, and Cos-10 metic Act (21 U.S.C. 321(g)(1)), biological product (as 11 defined in section 351(i) of the Public Health Service Act (42 U.S.C. 262(i))), or device (as defined in section 12 13 201(h) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 321(h))). 14