

MOTION TO RECOMMIT H.R. 5247

M____. _____ moves to recommit the bill H.R. 5247 to the Committee on Energy and Commerce with instructions to report the same back to the House forthwith, with the following amendment:

Strike section 2 and insert the following:

1 **SEC. 2. USE OF UNAPPROVED INVESTIGATIONAL DRUGS BY**
2 **PATIENTS DIAGNOSED WITH A TERMINAL**
3 **ILLNESS.**

4 (a) IN GENERAL.—Chapter V of the Federal Food,
5 Drug, and Cosmetic Act is amended by inserting after sec-
6 tion 561A (21 U.S.C. 360bbb–0) the following:

7 **“SEC. 561B. INVESTIGATIONAL DRUGS FOR USE BY ELIGI-**
8 **BLE PATIENTS.**

9 “(a) USE OF CLINICAL OUTCOMES.—

10 “(1) IN GENERAL.—The Secretary shall issue
11 guidance describing the Secretary’s consideration
12 and evaluation, for purposes of the review of, and
13 decision on whether to approve, a marketing applica-
14 tion under section 505 of this Act or section 351 of
15 the Public Health Service Act for an investigational
16 drug, of clinical outcomes associated with the provi-

1 sion by a sponsor or manufacturer of such drug
2 under subsection (b) or (c) of section 561. Such
3 guidance shall address—

4 “(A) specific instances in which the Sec-
5 retary will determine that the public health re-
6 quires such consideration and evaluation;

7 “(B) specific instances in which a sponsor
8 may request such consideration and evaluation;
9 and

10 “(C) the context in which such consider-
11 ation and evaluation will occur, particularly
12 with regard to information and data relevant to
13 the evaluation of a marketing application under
14 section 505 of this Act or section 351 of the
15 Public Health Service Act for the investiga-
16 tional drug.

17 “(2) GUIDANCE.—

18 “(A) DRAFT GUIDANCE.—Not later than 1
19 year after the date of enactment of this section,
20 the Secretary shall issue draft guidance with a
21 public comment period regarding the use of
22 clinical outcomes associated with the use of an
23 investigational drug that a sponsor or manufac-
24 turer has provided under subsection (b) or (c)
25 of section 561, as described in paragraph (1).

1 “(B) FINAL GUIDANCE.—Not later than 1
2 year after the public comment period on such
3 draft guidance ends, the Secretary shall issue
4 final guidance.

5 “(b) POSTING OF INFORMATION.—Not later than 1
6 year after the date of enactment of this section, the Sec-
7 retary shall post on the internet website of the Food and
8 Drug Administration and update annually, categorized by
9 therapeutic area—

10 “(1) the number of requests that were received
11 by the Food and Drug Administration for the provi-
12 sion by a sponsor or manufacturer of an investiga-
13 tional drug under subsection (b) or (c) of section
14 561; and

15 “(2) the number of such requests that were
16 granted.”.

17 (b) REPORTING.—Section 561A of the Federal Food,
18 Drug, and Cosmetic Act (21 U.S.C. 360bbb-0) is amend-
19 ed adding at the end the following:

20 “(g) REPORTING.—The manufacturer or sponsor of
21 an eligible investigational drug shall post on the same pub-
22 licly available internet website used by the manufacturer
23 for purposes of subsection (b) of this section an annual
24 summary of any provision by the manufacturer or sponsor
25 of an investigational drug under subsection (b) or (c) of

1 section 561. The summary shall include the number of re-
2 quests received, the number of requests granted, the num-
3 ber of patients treated, the therapeutic area of the drug
4 made available, and any known or suspected serious ad-
5 verse events. Such annual summary shall be provided to
6 the Secretary upon request.”.

7 (c) LIABILITY.—Section 561 of the Federal Food,
8 Drug, and Cosmetic Act (21 U.S.C. 360bbb) is amend-
9 ed—

10 (1) by redesignating subsection (e) as sub-
11 section (f); and

12 (2) by inserting after subsection (d) the fol-
13 lowing:

14 “(e) LIABILITY.—

15 “(1) ALLEGED ACTS OR OMISSIONS.—

16 “(A) MANUFACTURER OR SPONSOR.—No
17 manufacturer or sponsor (or their agent or rep-
18 resentative) of an investigational drug provided
19 to a single patient or small group of patients
20 for treatment use shall be liable for any alleged
21 act or omission related to the provision of such
22 drug, so long as such drug was provided in ac-
23 cordance with subsection (b) or (c), including
24 the reporting of safety information, from clin-
25 ical trials or any other source, as required pur-

1 suant to section 312.32 of title 21, Code of
2 Federal Regulations (or any successor regula-
3 tions).

4 “(B) PHYSICIAN, CLINICAL INVESTIGATOR,
5 OR HOSPITAL.—

6 “(i) No licensed physician, clinical in-
7 vestigator, or hospital shall be liable for
8 any alleged act or omission related to the
9 provision to a single patient or small group
10 of patients for treatment use of an inves-
11 tigational drug in accordance with the re-
12 quirements described in clause (ii), unless
13 such act or omission constitutes on the
14 part of such physician, clinical investigator,
15 or hospital with respect to such investiga-
16 tional drug—

17 “(I) willful or criminal mis-
18 conduct;

19 “(II) reckless misconduct;

20 “(III) gross negligence relative to
21 the applicable standard of care and
22 practice with respect to the adminis-
23 tration or dispensing of such inves-
24 tigational drug; or

1 “(IV) an intentional tort under
2 applicable State law.

3 “(ii) The requirements described in
4 this clause are the requirements under
5 subsection (b) or (c), including—

6 “(I) the reporting of safety infor-
7 mation, from clinical trials or any
8 other source, as required pursuant to
9 under section 312.32 of title 21, Code
10 of Federal Regulations (or any suc-
11 cessor regulations);

12 “(II) ensuring that the informed
13 consent requirements of part 50 of
14 title 21, Code of the Federal Regula-
15 tions (or any successor regulations)
16 are met; and

17 “(III) ensuring that review by an
18 institutional review board is obtained
19 in a manner consistent with the re-
20 quirements of part 56 of title 21,
21 Code of the Federal Regulations (or
22 any successor regulations).

23 “(2) DETERMINATION NOT TO PROVIDE
24 DRUG.—No manufacturer, sponsor, licensed physi-
25 cian, clinical investigator, or hospital, nor the Sec-

1 retary, shall be liable for determining not to provide
2 access to an investigational drug under this section
3 or for discontinuing any such access that it initially
4 determined to provide.

5 “(3) LIMITATION.—

6 “(A) IN GENERAL.—Except as set forth in
7 paragraphs (1) and (2), nothing in this section
8 or section 561B shall be construed to modify or
9 otherwise affect the right of any person to bring
10 a private action against a manufacturer or
11 sponsor (or their agent or representative), phy-
12 sician, clinical investigator, hospital, prescriber,
13 dispenser, or other entity under any State or
14 Federal product liability, tort, consumer protec-
15 tion, or warranty law.

16 “(B) FEDERAL GOVERNMENT.—Nothing in
17 this section or section 561B shall be construed
18 to modify or otherwise affect the authority of
19 the Federal Government to bring suit under
20 any Federal law.”.

