AMENDMENT TO RULES COMMITTEE PRINT 118-36

OFFERED BY MS. DEAN OF PENNSYLVANIA

Add at the end of subtitle C of title XVII the following:

1	SEC. 17 COMMENTED ACCELTED MEDICAL USE WITH
2	SEVERE RESTRICTIONS.
3	(a) Definitions.—Section 102 of the Controlled
4	Substances Act (21 U.S.C. 802) is amended—
5	(1) by redesignating paragraph (58) (defining a
6	serious violent felony) as paragraph (59);
7	(2) by redesignating the second paragraph (57)
8	(defining a serious drug felony) as paragraph (58);
9	and
10	(3) by adding at the end the following:
11	"(60)(A) Subject to subparagraph (B), the
12	term 'currently accepted medical use with severe re-
13	strictions', with respect to a drug or other sub-
14	stance, includes a drug or other substance that is an
15	active metabolite, moiety, or ingredient (whether in
16	natural or synthetic form) of an investigational new
17	drug for which a waiver is in effect under section
18	505(i) of the Federal Food, Drug, and Cosmetic Act

1	(21 U.S.C. 355(i)) or section 351(a)(3) of the Public
2	Health Service Act (42 U.S.C. 262(a)(3)) and that
3	the Secretary—
4	"(i) designates as a breakthrough therapy
5	under section 506(a) of the Federal Food,
6	Drug, and Cosmetic Act (21 U.S.C. 356(a)); or
7	"(ii) authorizes for expanded access under
8	subsection (b) or (c) of section 561 of the Fed-
9	eral Food, Drug, and Cosmetic Act (21 U.S.C.
10	360bbb), either alone or as part of a thera-
11	peutic protocol, to treat patients with serious or
12	life-threatening diseases for which no com-
13	parable or satisfactory therapies are available.
14	"(B) A drug or other substance shall not be
15	treated as meeting the criteria under subparagraph
16	(A) for having a currently accepted medical use with
17	severe restrictions if—
18	"(i) in the case of a drug or other sub-
19	stance described in subparagraph (A)(ii)—
20	"(I) the Secretary places the ex-
21	panded access or protocol for such drug on
22	clinical hold as described in section 312.42
23	of title 21, Code of Federal Regulations (or
24	any successor regulations);

1	"(II) there is no other investigational
2	new drug containing the drug or other sub-
3	stance for which expanded access has been
4	authorized under section 561(a) of the
5	Federal Food, Drug, and Cosmetic Act (21
6	U.S.C. 360bbb(a)); and
7	"(III) the drug or other substance
8	does not meet the requirements of sub-
9	paragraph (A)(i); or
10	"(ii) the drug or other substance is ap-
11	proved under section 505 of the Federal Food,
12	Drug, and Cosmetic Act (21 U.S.C. 355) or
13	section 351 of the Public Health Service Act
14	(42 U.S.C. 262).".
15	(b) AUTHORITY AND CRITERIA FOR CLASSIFICATION
16	OF SUBSTANCES.—Section 201(j) of the Controlled Sub-
17	stances Act (21 U.S.C. 811(j)) is amended—
18	(1) in paragraph (1), by inserting "a drug des-
19	ignated as a breakthrough therapy under section
20	506(a) of the Food Drug and Cosmetic Act (21
21	U.S.C. 356(a)), or a drug authorized for expanded
22	access under subsection (b) or (c) of section 561 of
23	the Federal Food, Drug, and Cosmetic Act (21
24	U.S.C. 360bbb)," after "subsection (f),";
25	(2) in paragraph (2)—

1	(A) in subparagraph (A), by striking ";
2	or" and inserting a semicolon;
3	(B) in subparagraph (B), by striking the
4	period at the end and inserting a semicolon;
5	and
6	(C) by adding at the end the following:
7	"(C) the date on which the Attorney Gen-
8	eral receives notification from the Secretary of
9	Health and Human Services that the Secretary
10	has designated the drug as a breakthrough
11	therapy under section 506(a) of the Federal
12	Food, Drug, and Cosmetic Act (21 U.S.C.
13	356(a)) or authorized the drug for expanded ac-
14	cess under subsection (b) or (c) of section 561
15	of the Federal Food, Drug, and Cosmetic Act
16	(21 U.S.C. 360bbb); or
17	"(D) the date on which the Attorney Gen-
18	eral receives any written notification dem-
19	onstrating that the Secretary, before the date of
20	enactment of this subparagraph, designated the
21	drug as a breakthrough therapy under section
22	506(a) of the Federal Food, Drug, and Cos-
23	metic Act (21 U.S.C. 356(a)) or authorized the
24	drug for expanded access under subsection (b)

1	or (c) of section 361 of the Federal Food,
2	Drug, and Cosmetic Act (21 U.S.C. 360bbb).";
3	(3) in paragraph (3), by inserting "or para-
4	graph (4)" after "paragraph (1)"; and
5	(4) by adding at the end the following:
6	"(4) With respect to a drug moved from schedule I
7	to schedule II pursuant to paragraph (1) and the expe-
8	dited procedures described under this subsection, if the
9	drug no longer has a currently accepted medical use with
10	severe restrictions and the Secretary of Health and
11	Human Services recommends that the Attorney General
12	control the drug in schedule I pursuant to subsections (a)
13	and (b), the Attorney General shall, not later than 90 days
14	after receiving written notification from the Secretary,
15	issue an interim final rule controlling the drug in accord-
16	ance with such subsections and section 202(b) using the
17	procedures described in paragraph (3) of this subsection.".

