

AMENDMENT IN THE NATURE OF A SUBSTITUTE
TO H.R. 1266
OFFERED BY M. _____

Strike all after the enacting clause and insert the following:

1 **SECTION 1. SHORT TITLE.**

2 This Act may be cited as the “Combating Illicit
3 Xylazine Act”.

4 SEC. 2. DEFINITIONS.

(a) IN GENERAL.—In this Act, the term “xylazine” has the meaning given the term in paragraph (61) of section 102 of the Controlled Substances Act, as added by subsection (b) of this section.

9 (b) CONTROLLED SUBSTANCES ACT.—Section 102 of
10 the Controlled Substances Act (21 U.S.C. 802) is amend-
11 ed by adding at the end the following:

“(61) The term ‘xylazine’ means the substance
xylazine, including its salts, isomers, and salts of isomers
whenever the existence of such salts, isomers, and salts
of isomers is possible.”.

1 **SEC. 3. ADDING XYLAZINE TO SCHEDULE III.**

2 Schedule III of section 202(c) of the Controlled Sub-
3 stances Act (21 U.S.C. 812) is amended by adding at the
4 end the following:

5 “(f) Unless specifically excepted or unless listed in
6 another schedule, any material, compound, mixture, or
7 preparation which contains any quantity of xylazine.”.

8 **SEC. 4. AMENDMENTS.**

9 (a) AMENDMENT.—Section 102 of the Controlled
10 Substances Act (21 U.S.C. 802) is amended by striking
11 paragraph (27) and inserting the following:

12 “(27)(A) Except as provided in subparagraph (B),
13 the term ‘ultimate user’ means a person who has lawfully
14 obtained, and who possesses, a controlled substance for
15 the use by the person or for the use of a member of the
16 household of the person or for an animal owned by the
17 person or by a member of the household of the person.

18 “(B)(i) In the case of xylazine, other than for a drug
19 product approved under subsection (b) or (j) of section
20 505 of the Federal Food, Drug, and Cosmetic Act (21
21 U.S.C. 355), the term ‘ultimate user’ means a person—

22 “(I) to whom xylazine was dispensed by—

23 “(aa) a veterinarian registered under this

24 Act; or

1 “(bb) a pharmacy registered under this
2 Act pursuant to a prescription of a veterinarian
3 registered under this Act; and

4 “(II) who possesses xylazine for—

5 “(aa) an animal owned by the person or by
6 a member of the household of the person;

7 “(bb) an animal under the care of the per-
8 son;

9 “(cc) use in government animal-control
10 programs authorized under applicable Federal,
11 State, Tribal, or local law; or

12 “(dd) use in wildlife programs authorized
13 under applicable Federal, State, Tribal, or local
14 law.

15 “(ii) In this subparagraph, the term ‘person’ in-
16 cludes—

17 “(I) a government agency or business where
18 animals are located; and

19 “(II) an employee or agent of an agency or
20 business acting within the scope of their employment
21 or agency.”.

22 (b) FACILITIES.—An entity that manufactures
23 xylazine, as of the date of enactment of this Act, shall
24 not be required to make capital expenditures necessary to
25 install the security standard required of schedule III of

1 the Controlled Substances Act (21 U.S.C. 801 et seq.) for
2 the purposes of manufacturing xylazine.

3 (c) LABELING.—The requirements related to label-
4 ing, packaging, and distribution logistics of a controlled
5 substance in schedule III of section 202(c) of the Con-
6 trolled Substances Act (21 U.S.C. 812(c)) shall not take
7 effect for xylazine until the date that is 1 year after the
8 date of enactment of this Act.

9 (d) PRACTITIONER REGISTRATION.—The require-
10 ments related to practitioner registration, inventory, and
11 recordkeeping of a controlled substance in schedule III of
12 section 202(c) of the Controlled Substances Act (21
13 U.S.C. 812(c)) shall not take effect for xylazine until the
14 date that is 60 days after the date of enactment of this
15 Act. A practitioner that has applied for registration during
16 the 60-day period beginning on the date of enactment of
17 this Act may continue their lawful activities until such ap-
18 plication is approved or denied.

19 (e) MANUFACTURER TRANSITION.—The Food and
20 Drug Administration and the Drug Enforcement Adminis-
21 tration shall facilitate and expedite the relevant manufac-
22 turer submissions or applications required by the place-
23 ment of xylazine on schedule III of section 202(c) of the
24 Controlled Substances Act (21 U.S.C. 812(c)).

1 (f) CLARIFICATION.—Nothing in this Act, or the
2 amendments made by this Act, shall be construed to re-
3 quire the registration of an ultimate user of xylazine under
4 the Controlled Substances Act (21 U.S.C. 801 et seq.) in
5 order to possess xylazine in accordance with subparagraph
6 (B) of section 102(27) of that Act (21 U.S.C. 802(27)),
7 as added by subsection (a) of this section.

8 **SEC. 5. ARCOS TRACKING.**

9 Section 307(i) of the Controlled Substances Act (21
10 U.S.C. 827(i)) is amended—

11 (1) in the matter preceding paragraph (1)—

12 (A) by inserting “or xylazine” after
13 “gamma hydroxybutyric acid”;

14 (B) by inserting “or 512” after “section
15 505”; and

16 (C) by inserting “respectively,” after “the
17 Federal Food, Drug, and Cosmetic Act,”; and

18 (2) in paragraph (6), by inserting “or xylazine”
19 after “gamma hydroxybutyric acid”.

20 **SEC. 6. SENTENCING COMMISSION.**

21 Pursuant to its authority under section 994(p) of title
22 28, United States Code, the United States Sentencing
23 Commission shall review and, if appropriate, amend its
24 sentencing guidelines, policy statements, and official com-
25 mentary applicable to persons convicted of an offense

1 under section 401 of the Controlled Substances Act (21
2 U.S.C. 841) or section 1010 of the Controlled Substances
3 Import and Export Act (21 U.S.C. 960) to provide appro-
4 priate penalties for offenses involving xylazine that are
5 consistent with the amendments made by this Act. In car-
6 rying out this section, the Commission should consider the
7 common forms of xylazine as well as its use alongside
8 other scheduled substances.

9 **SEC. 7. REPORT TO CONGRESS ON XYLAZINE.**

10 (a) INITIAL REPORT.—Not later than 18 months
11 after the date of the enactment of this Act, the Attorney
12 General, acting through the Administrator of the Drug
13 Enforcement Administration and in coordination with the
14 Commissioner of Food and Drugs, shall submit to Con-
15 gress a report on the prevalence of illicit use of xylazine
16 in the United States and the impacts of such use, includ-
17 ing—

- 18 (1) where the drug is being diverted;
19 (2) where the drug is originating; and
20 (3) whether any analogues to xylazine, or re-
21 lated or derivative substances, exist and present a
22 substantial risk of abuse.

23 (b) ADDITIONAL REPORT.—Not later than 4 years
24 after the date of the enactment of this Act, the Attorney
25 General, acting through the Administrator of the Drug

1 Enforcement Administration and in coordination with the
2 Commissioner of Food and Drugs, shall submit to Con-
3 gress a report updating Congress on the prevalence and
4 proliferation of xylazine trafficking and misuse in the
5 United States.

