118TH CONGRESS 1ST SESSION				S	S.			

To direct the Secretary of Veterans Affairs to carry out a study and clinical trials on the effects of cannabis on certain health outcomes of veterans with chronic pain and post-traumatic stress disorder, and for other purposes.

IN THE SENATE OF THE UNITED STATES

Mr.	TESTER	introduced	the following	; bill;	which	was	read	twice	and	referre	d
		to the Co	ommittee on								

A BILL

- To direct the Secretary of Veterans Affairs to carry out a study and clinical trials on the effects of cannabis on certain health outcomes of veterans with chronic pain and post-traumatic stress disorder, 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 "VA Medicinal Cannabis
 - 5 Research Act of 2023".
 - 6 SEC. 2. DEFINITIONS.
 - 7 In this Act:

1	(1) COVERED VETERAN.—The term "covered
2	veteran" means a veteran who is enrolled in the pa-
3	tient enrollment system of the Department of Vet-
4	erans Affairs established and operated under section
5	1705(a) of title 38, United States Code.
6	(2) Secretary.— The term "Secretary"
7	means the Secretary of Veterans Affairs.
8	SEC. 3. DEPARTMENT OF VETERANS AFFAIRS LARGE-
9	SCALE, MIXED METHODS, RETROSPECTIVE
10	QUALITATIVE STUDY ON THE EFFECTS OF
11	CANNABIS ON CERTAIN HEALTH OUTCOMES
12	OF VETERANS WITH CHRONIC PAIN AND
13	POST-TRAUMATIC STRESS DISORDER.
13 14	POST-TRAUMATIC STRESS DISORDER. (a) Study Required.—
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14 15 16 17 18	(a) Study Required.— (1) In General.—The Secretary, through the Office of Research and Development of the Department of Veterans Affairs, shall carry out a large-scale, mixed methods, retrospective, and qualitative
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14 15 16 17 18 19 20	(a) Study Required.— (1) In General.—The Secretary, through the Office of Research and Development of the Department of Veterans Affairs, shall carry out a large-scale, mixed methods, retrospective, and qualitative study on the effects of cannabis on the health outcomes of covered veterans diagnosed with chronic
14 15 16 17 18 19 20 21	(a) Study Required.— (1) In General.—The Secretary, through the Office of Research and Development of the Department of Veterans Affairs, shall carry out a large-scale, mixed methods, retrospective, and qualitative study on the effects of cannabis on the health outcomes of covered veterans diagnosed with chronic pain and covered veterans diagnosed with post-trau-

1	observational study on the effects of cannabis use on
2	the health of covered veterans.
3	(3) Elements.—
4	(A) In general.—The study required by
5	paragraph (1) shall—
6	(i) triangulate a range of data
7	sources;
8	(ii) compare the positive and negative
9	health outcomes of covered veterans who
10	use cannabis, utilizing outcomes that can
11	be measured in an electronic health record
12	of the Department and through data sets
13	of the Department relating to claims for
14	benefits under the laws administered by
15	the Secretary;
16	(iii) elicit the positive and negative
17	outcomes of cannabis use for covered vet-
18	erans through semi-structured interviews;
19	(iv) estimate current and future
20	health system needs to address positive
21	and negative outcomes of cannabis use for
22	covered veterans;
23	(v) include a qualitative, open-ended
24	survey provided to covered veterans who
25	have sought care from the Department for

1	chronic pain or post-traumatic stress dis-
2	order during the five-year period preceding
3	the survey; and
4	(vi) include an assessment of—
5	(I) all records within the Vet-
6	erans Health Administration for cov-
7	ered veterans participating in the
8	study; and
9	(II) all records within the Vet-
10	erans Benefits Administration for cov-
11	ered veterans participating in the
12	study.
13	(B) Health outcomes.—A comparison
14	of health outcomes under subparagraph (A)(ii)
15	shall include an assessment of the following:
16	(i) The reduction or increase in opiate
17	use or dosage.
18	(ii) The reduction or increase in
19	benzodiazepine use or dosage.
20	(iii) The reduction or change in use of
21	other types of medication.
22	(iv) The reduction or increase in alco-
23	hol use.
24	(v) The reduction or increase in the
25	prevalence of substance abuse disorders.

1	(vi) Sleep quality.
2	(vii) Osteopathic pain (including pain
3	intensity and pain-related outcomes).
4	(viii) Agitation.
5	(ix) Quality of life.
6	(x) Mortality and morbidity.
7	(xi) Hospital readmissions.
8	(xii) Any newly developed or exacer-
9	bated health conditions, including mental
10	health conditions.
11	(b) Implementation.—Not later than 180 days
12	after the date of the enactment of this Act, the Secretary
13	shall commence the implementation of the study required
14	by subsection $(a)(1)$.
15	(c) Duration of Study.—The study required by
16	subsection (a)(1) shall be carried out for an 18-month pe-
17	riod.
18	(d) Report.—
19	(1) In general.—Not later than 90 days after
20	the completion of the study required by subsection
21	(a)(1), the Secretary shall submit to the Committee
22	on Veterans' Affairs of the Senate and the Com-
23	mittee on Veterans' Affairs of the House of Rep-
24	resentatives a report on the study.

1 (2) Ability to conduct clinical trials.— 2 The Secretary shall include in the report required by 3 paragraph (1) an assessment of whether the Sec-4 retary is able to meet the criteria necessary to con-5 duct the clinical trials required under section 4, in-6 cluding consideration of subsection (e)(1) of such 7 section. 8 SEC. 4. DEPARTMENT OF VETERANS AFFAIRS CLINICAL 9 TRIALS ON THE EFFECTS OF CANNABIS ON 10 CERTAIN HEALTH OUTCOMES OF VETERANS 11 WITH CHRONIC PAIN AND POST-TRAUMATIC 12 STRESS DISORDER. 13 (a) CLINICAL TRIALS REQUIRED.— 14 (1) IN GENERAL.—If the Secretary indicates in 15 the report required by section 3(d) that the Sec-16 retary is able to meet the criteria necessary to pro-17 ceed to clinical trials, commencing not later than 18 180 days after the submittal of that report, the Sec-19 retary shall carry out a series of clinical trials on the 20 effects of cannabis appropriate for investigational 21 use, as determined by the Food and Drug Adminis-22 tration under section 505(i) of the Federal Food, 23 Drug, and Cosmetic Act (21 U.S.C. 355(i)), on the 24 health outcomes of covered veterans diagnosed with

1	chronic pain and covered veterans diagnosed with
2	post-traumatic stress disorder.
3	(2) Considerations.—The clinical trials re-
4	quired by paragraph (1) shall include, as appro-
5	priate, an evaluation of key symptoms, clinical out-
6	comes, and conditions associated with chronic pain
7	and post-traumatic stress disorder, which may in-
8	clude—
9	(A) with respect to covered veterans diag-
10	nosed with chronic pain, an evaluation of the
11	effects of the use of cannabis on—
12	(i) osteopathic pain (including pain in-
13	tensity and pain-related outcomes);
14	(ii) the reduction or increase in opioid
15	use or dosage;
16	(iii) the reduction or increase in
17	benzodiazepine use or dosage;
18	(iv) the reduction or increase in alco-
19	hol use;
20	(v) the reduction or increase in the
21	prevalence of substance use disorders;
22	(vi) inflammation;
23	(vii) sleep quality;
24	(viii) agitation;
25	(ix) quality of life;

1	(x) exacerbated or new mental health
2	conditions; and
3	(xi) suicidal ideation.
4	(B) with respect to covered veterans diag-
5	nosed with post-traumatic stress disorder, an
6	evaluation of the effects of the use of cannabis
7	on—
8	(i) the symptoms of post-traumatic
9	stress disorder (PTSD) as established by
10	or derived from the clinician administered
11	PTSD scale, the PTSD checklist, the
12	PTSD symptom scale, the post-traumatic
13	diagnostic scale, and other applicable
14	methods of evaluating symptoms of post-
15	traumatic stress disorder;
16	(ii) the reduction or increase in
17	benzodiazepine use or dosage;
18	(iii) the reduction or increase in alco-
19	hol use;
20	(iv) the reduction or increase in the
21	prevalence of substance use disorders;
22	(v) mood;
23	(vi) anxiety;
24	(vii) social functioning;
25	(viii) agitation;

1	(ix) suicidal ideation; and
2	(x) sleep quality, including frequency
3	of nightmares and night terrors.
4	(3) Optional elements.—The clinical trials
5	required by paragraph (1) may include, as appro-
6	priate, an evaluation of the effects of the use of can-
7	nabis to treat chronic pain and post-traumatic stress
8	disorder on other symptoms, clinical outcomes, and
9	conditions not covered by paragraph (2), which may
10	include—
11	(A) pulmonary function;
12	(B) cardiovascular events;
13	(C) head, neck, and oral cancer;
14	(D) testicular cancer;
15	(E) ovarian cancer;
16	(F) transitional cell cancer;
17	(G) intestinal inflammation;
18	(H) motor vehicle accidents; or
19	(I) spasticity.
20	(b) Long-term Observational Study.—The Sec-
21	retary may carry out a long-term observational study of
22	the participants in the clinical trials required by sub-
23	section (a).
24	(c) Type of Cannabis.—

1	(1) In general.—In carrying out the clinical
2	trials required by subsection (a), the Secretary shall
3	study varying forms of cannabis, including whole
4	plant raw material and extracts, and may study
5	varying routes of administration.
6	(2) Plant cultivars.—Of the varying forms
7	of cannabis required under paragraph (1), the Sec-
8	retary shall study plant cultivars with varying ratios
9	of tetrahydrocannabinol to cannabidiol.
10	(d) Implementation.—Not later than 18 months
11	after the date of the enactment of this Act, the Secretary
12	shall—
13	(1) develop a plan to implement this section
14	and submit such plan to the Committee on Veterans
15	Affairs of the Senate and the Committee on Vet-
16	erans' Affairs of the House of Representatives; and
17	(2) issue any requests for proposals the Sec-
18	retary determines appropriate for such implementa-
19	tion.
20	(e) TERMINATION OF CLINICAL TRIALS.—
21	(1) CLINICAL GUIDELINE REQUIREMENTS OF
22	EXCESSIVE RISK.—The Secretary may terminate the
23	clinical trials required by subsection (a) if the Sec-
24	retary determines that the Department of Veterans
25	Affairs is unable to meet clinical guideline require-

ments necessary to conduct such trials or the clinical
trials would create excessive risk to participants.

(2) Completion upon submittal of final report required under subsection (f)(2).

(f) Reports.—

- (1) Periodic Reports.—During the five-year period beginning on the date of the commencement of clinical trials required by subsection (a), the Secretary shall submit periodically, but not less frequently than annually, to the Committee on Veterans' Affairs of the Senate and the Committee on Veterans' Affairs of the House of Representatives reports on the implementation of this section.
- (2) Final Report.—Not later than one year after the completion of the five-year period specified in paragraph (1), the Secretary shall submit to the Committee on Veterans' Affairs of the Senate and the Committee on Veterans' Affairs of the House of Representatives a final report on the implementation of this section.

23 SEC. 5. ADMINISTRATION OF STUDY AND CLINICAL TRIALS.

24 (a) Demographic Representation.—In carrying 25 out the study required by section 3 and the clinical trials

- 1 required by section 4, the Secretary shall ensure represen-
- 2 tation in such study and trials of demographics that rep-
- 3 resent the population of veterans in the United States, as
- 4 determined by the most recently available data from the
- 5 American Community Survey of the Bureau of the Census.
- 6 (b) Data Preservation.—The Secretary shall en-
- 7 sure that the study required by section 3 and the clinical
- 8 trials required by section 4 include a mechanism to en-
- 9 sure—
- 10 (1) the preservation of all data, including all
- data sets and survey results, collected or used for
- purposes of such study and trials in a manner that
- will facilitate further research; and
- 14 (2) registration of such data in the database of
- privately and publicly funded clinical studies main-
- tained by the National Library of Medicine (or suc-
- 17 cessor database).
- 18 (c) Anonymous Data.—The Secretary shall ensure
- 19 that data relating to any study or clinical trial conducted
- 20 under this Act is anonymized and cannot be traced back
- 21 to an individual patient.
- 22 (d) Effect on Other Benefits.—The eligibility
- 23 or entitlement of a covered veteran to any other benefit
- 24 under the laws administered by the Secretary or any other
- 25 provision of law shall not be affected by the participation

1	of the covered veteran in the study under section 3, a clin-
2	ical trial under section 4(a), or a study under section 4(b)
3	(e) Effect on Other Laws.—Nothing in this Act
4	shall affect or modify—
5	(1) the Federal Food, Drug, and Cosmetic Act
6	(21 U.S.C. 301 et seq.);
7	(2) section 351 of the Public Health Service
8	Act (42 U.S.C. 262); or
9	(3) the authority of the Commissioner of Food
10	and Drugs and the Secretary of Health and Human
11	Services—
12	(A) under—
13	(i) the Federal Food, Drug, and Cos-
14	metic Act (21 U.S.C. 301 et seq.); or
15	(ii) section 351 of the Public Health
16	Service Act (42 U.S.C. 262); or
17	(B) to promulgate Federal regulations and
18	guidelines pertaining to cannabidiol, marijuana
10	or other subject matter addressed in this Act