110TH CONGRESS 1ST SESSION H.R. 2717

To permit an individual to be treated by a health care practitioner with any method of medical treatment such individual requests, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

JUNE 14, 2007

Mr. BURTON of Indiana introduced the following bill; which was referred to the Committee on Energy and Commerce

A BILL

- To permit an individual to be treated by a health care practitioner with any method of medical treatment such individual requests, 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 "Access to Medical

5 Treatment Act".

6 SEC. 2. DEFINITIONS.

- 7 In this Act:
- 8 (1) ADVERTISING CLAIM.—The term "adver9 tising claim" means any representation made or sug-

1	gested by statement, word, design, device, sound, or
2	any combination thereof with respect to a medical
3	treatment.
4	(2) DANGER.—The term "danger" means an
5	adverse reaction to an unapproved drug or medical
6	device that, when used as directed—
7	(A) causes serious harm;
8	(B) occurred as a result of the medical
9	treatment;
10	(C) would not otherwise have occurred;
11	and
12	(D) is more serious than reactions experi-
13	enced with routinely used medical treatments
14	approved by the Food and Drug Administration
15	for the same medical condition or conditions.
16	(3) DEVICE.—The term "device" has the mean-
17	ing given such term in section 201(h) of the Federal
18	Food, Drug, and Cosmetic Act (21 U.S.C.321(h)).
19	(4) Drug.—The term "drug" has the meaning
20	given such term in section $201(g)(1)$ of the Federal
21	Food, Drug, and Cosmetic Act (21 U.S.C. 321
22	(g)(1)).
23	(5) FOOD.—The term "food"—

1	(A) has the meaning given such term in
2	section 201(f) of the Federal Food, Drug, and
3	Cosmetic Act (21 U.S.C. 321(f)); and
4	(B) includes a dietary supplement as de-
5	fined in section 201(ff) of such Act.
6	(6) HEALTH CARE PRACTITIONER.—The term
7	"health care practitioner" means a physician or
8	other individual who is legally authorized to provide
9	health care services in the State in which the serv-
10	ices are provided.
11	(7) INTERSTATE COMMERCE.—The term "inter-
12	state commerce'' means commerce between any
13	State or territory and any place outside thereof, and
14	commerce within the District of Columbia or within
15	any other territory not organized with a legislative
16	body.
17	(8) LABEL.—The term "label" has the meaning
18	given such term in section 201(k) of the Federal
19	Food, Drug, and Cosmetic Act (21 U.S.C. 321(k)).
20	(9) LABELING.—The term "labeling" has the
21	meaning given such term in section 201(m) of the
22	Federal Food, Drug, and Cosmetic Act (21 U.S.C.
23	321(m)).
24	(10) LEGAL REPRESENTATIVE.—The term

25 "legal representative" means a parent or an indi-

vidual who qualifies as a legal guardian under appli cable State law.

3 (11) MEDICAL DEVICE.—The term "medical de4 vice" has the meaning given the term "device" in
5 section 201(h) of the Federal Food, Drug, and Cos6 metic Act (21 U.S.C. 321(h)).

7 (12) MEDICAL TREATMENT.—The term "med8 ical treatment" means any food, drug, device, or
9 procedure that is used and intended as a cure, miti10 gation, treatment, or prevention of disease or a
11 health condition.

(13) PATIENT.—The term "patient" means any
individual who seeks medical treatment from a
health care practitioner for a disease or health condition.

SECRETARY.—The 16 (14)term "Secretary" 17 means the Secretary of Health and Human Services. (15) Seller.—The term "seller" means an in-18 19 dividual or organization that receives payment re-20 lated to the medical treatment of a patient of a 21 health practitioner, except that this term does not 22 apply to a health care practitioner who receives pay-23 ment from an individual or representative of such in-24 dividual for the administration of a medical treat-25 ment to such individual.

1 (16) UNAPPROVED DRUG OR MEDICAL DE-2 VICE.—The term "unapproved drug or medical device" with respect to a drug or medical device, 3 4 means a drug or medical device that is not approved 5 or authorized for manufacture, sale, and distribution 6 in interstate commerce under section 505, 510, or 7 515 of the Federal Food, Drug, and Cosmetic Act 8 (21 U.S.C 355, 360c, and 360(e)) or under section 9 351 of the Public Health Service Act (42 U.S.C. 10 262).

11 SEC. 3. ACCESS TO MEDICAL TREATMENT.

12 (a) IN GENERAL.—Notwithstanding any other provi-13 sion of law, and except as provided in subsection (b), an individual shall have the right to be treated by a health 14 15 care practitioner with any medical treatment (including a medical treatment that is not approved, certified, or li-16 17 censed by the Secretary) that such individual desires, or that the legal representative of such individual authorizes, 18 19 if—

20 (1) such practitioner has personally examined
21 such individual and agrees to provide treatment to
22 such individual;

23 (2) the administration of such treatment does
24 not violate applicable licensing laws;

1 (3) the health care practitioner complies with 2 the requirements of subsection (b); and 3 (4) it is a medical treatment that has not been 4 approved, certified, or licensed by the Secretary, or 5 is any medical treatment that has been approved by 6 the designated governmental agency for a member 7 country of the European Union or the European 8 Free Trade Association, Canada, Australia, New 9 Zealand, or Japan but not otherwise approved, cer-

10 tified, or licensed by the Secretary.

11 (b) MEDICAL TREATMENT REQUIREMENTS.—

12 (1) IN GENERAL.—A health care practitioner
13 may provide the medical treatment requested by an
14 individual described in subsection (a) if—

(A) there is no reason for the practitioner
to conclude that, based on generally accepted
principles and current information, the medical
treatment requested, when used or provided as
directed, will cause danger to the patient;

(B) in the case of an individual whose
treatment is the administration of a food, drug,
or device that has to be approved, certified, or
licensed by the Secretary, but has not been so
approved, certified, or licensed—

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1	(i) such individual has been informed
2	in writing that such food, drug, or device
3	has not been approved, certified, or li-
4	censed by the Secretary for use as a med-
5	ical treatment of the medical condition of
6	such individual; and
7	(ii) prior to the administration of such
8	treatment, the practitioner has provided
9	the patient a written statement that in-
10	cludes the following provision: "WARN-
11	ING: This food, drug, or device has not
12	been declared to be safe and effective by
13	the Federal Government and any indi-
14	vidual who uses such food, drug, or device
15	does so at his or her own risk.";
16	(C) such individual has been informed in
17	writing of the nature of the medical treatment,
18	including-
19	(i) the contents and methods of such
20	treatment;
21	(ii) the anticipated benefits of such
22	treatment;
23	(iii) any reasonably foreseeable side
24	effects that may result from such treat-
25	ment;

1	(iv) the results of past application of
2	such treatment by the health care practi-
3	tioner and others; and
4	(v) any other information necessary to
5	fully meet the requirements for informed
6	consent of human subjects prescribed by
7	regulations issued by the Food and Drug
8	Administration;
9	(D) except as provided in subsection (c),
10	there have been no advertising claims made
11	with respect to the efficacy of the medical treat-
12	ment by the practitioner, manufacturer, or dis-
13	tributor;
14	(E) the label or labeling of any food, drug,
15	or device that is a part of the requested medical
16	treatment is not false or misleading;
17	(F) such individual—
18	(i) has been provided with a written
19	statement that such individual has been
20	fully informed with respect to the informa-
21	tion described in subparagraphs (A)
22	through (D);
23	(ii) desires such treatment; and
24	(iii) signs such statement; and

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1 (G) the health care practitioner provides 2 the patient with a recommendation for the treatment involved under circumstances that 3 4 give the patient sufficient opportunity to con-5 sider whether or not to use such treatment. 6 (2) BURDEN OF PROOF.—In any proceeding re-7 lating to the enforcement of paragraph (1)(E) with 8 respect to the label of a drug, device, or food used 9 in medical treatment covered under this subsection, 10 the provisions of section 403B(c) of the Federal 11 Food, Drug, and Cosmetic Act (21 U.S.C. 343–2(c)) 12 shall apply with respect to establishing the burden of 13 proof that such label is false or misleading. 14 (3) RULE OF CONSTRUCTION.—Nothing in this 15 section shall be construed to require informed con-16 sent for the prescription of dietary supplements and 17 foods not requiring such informed consent prior to 18 the date of the enactment of this Act. 19 (c) CLAIM EXCEPTIONS.— 20 (1) Reporting by a health care practi-21 TIONER.—Subsection (b)(1)(D) shall not apply to an 22 accurate and truthful reporting by a health care 23 practitioner of the results of the practitioner's ad-24 ministration of a medical treatment in recognized

25 journals, at seminars, conventions, or similar meet-

1 ings, or to others, so long as the reporting practi-2 tioner has no direct or indirect financial interest in 3 the reporting of the material and has received no fi-4 nancial benefits of any kind from the manufacturer, 5 distributor, or other seller for such reporting. Such 6 reporting may not be used by a manufacturer, dis-7 tributor, or other seller to advance the sale of such 8 treatment.

9 (2) STATEMENTS BY A PRACTITIONER TO A PA-10 TIENT.—Subsection (b)(1)(D) shall not apply to any 11 statement made by a health care practitioner di-12 rectly to a patient or prospective patient. A health 13 care practitioner shall not be held liable for any ad-14 vertising claims made by others unless the practi-15 tioner is a party in the dissemination of the informa-16 tion in such claims.

17 (3) DIETARY SUPPLEMENTS STATEMENT.—
18 Subsection (b)(1)(D) shall not apply to statements
19 or claims permitted under sections 403B and
20 403(r)(6) of the Federal Food, Drug, and Cosmetic
21 Act (21 U.S.C. 343–2 and 343(r)(6)).

22 SEC. 4. REPORTING OF A DANGEROUS MEDICAL TREAT23 MENT.

24 (a) HEALTH CARE PRACTITIONER.—If a health care25 practitioner, after administering a medical treatment, dis-

1	covers that the treatment itself was a danger to the indi-
2	vidual receiving such treatment, the practitioner shall—
3	(1) immediately cease the use of such treat-
4	ment;
5	(2) refrain from recommending the use of any
6	unapproved drug or medical device that was a part
7	of such treatment;
8	(3) report to the manufacturer and the Director
9	of the Centers for Disease Control and Prevention—
10	(A) the nature of such treatment;
11	(B) the results of such treatment;
12	(C) the complete protocol of such treat-
13	ment; and
14	(D) the source from which such treatment
15	or any part thereof was obtained; and
16	(4) include as part of the reporting under para-
17	graph (3) , an affidavit pursuant to section 1746 of
18	title 28, United States Code, confirming that all
19	statements made in the report under such paragraph
20	are accurate.
21	(b) Secretary.—Upon confirmation that a medical
22	treatment has proven dangerous to individuals, the Sec-
23	retary shall properly disseminate information with respect
24	to the danger of the medical treatment and prohibit the
25	further use of such treatment.

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3 If a health care practitioner, after administering a medical treatment that is not an approved drug or medical 4 5 device for a life-threatening medical condition or conditions, discovers that such medical treatment has, in the 6 7 opinion of the health care practitioner, positive effects on 8 such condition or conditions that are significantly greater 9 than the positive effects that are expected from an approved medical treatment for the same condition or condi-10 11 tions, the practitioner shall—

(1) make a monthly reporting to the National
Center for Complementary and Alternative Medicine
at the National Institutes of Health of—

15 (A) the nature of such medical treatment 16 (which is not a conventional medical treatment); 17 (B) the general results of such treatment 18 administered in the month involved; and 19 (C) the protocol of such treatment; and 20 (2) provide an affidavit pursuant to section 21 1746 of title 28, United States Code, confirming 22 that all statements made in the monthly reporting

23 under paragraph (1) are accurate and truthful.

1SEC. 6. TRANSPORTATION AND PRODUCTION OF FOOD,2DRUGS, DEVICES, AND OTHER EQUIPMENT.

3 (a) IN GENERAL.—Notwithstanding any other provi4 sion of the Federal Food, Drug, and Cosmetic Act (21)
5 U.S.C. 201 et seq.), an individual may—

6 (1) introduce or deliver into interstate com7 merce a food, drug, device, or any other equipment;
8 and

9 (2) produce, transport, receive and hold a food,
10 drug, device, or any other equipment,

11 solely for use in accordance with this Act if there have12 been no advertising claims by the manufacturer, dis-13 tributor, or seller of the food, drug, device, or equipment14 involved.

15 (b) RULE OF CONSTRUCTION.—Nothing in this Act 16 shall be construed to limit or interfere with the authority 17 of a health care practitioner to prescribe, recommend, pro-18 vide, or administer to a patient for any medical condition 19 or disease any unapproved drug or medical device that is 20 lawful under the law of the State or States in which the 21 health care practitioner practices.

22 SEC. 7. OTHER LAWS NOT AFFECTED BY THIS ACT.

23 Nothing in this Act shall be construed to—

(1) apply to the manufacturer, distribution,possession, or use of any drug that is a controlled

1	substance under the Controlled Substances Act $(21$
2	U.S.C. 801 et seq.);
3	(2) apply to statements or claims permitted or
4	authorized under sections 403 and 403B of the Fed-
5	eral Food, Drug, and Cosmetic Act (21 U.S.C. 343,
6	343–2); or
7	(3) in any way adversely affect the distribution
8	or sale of dietary supplements (as defined in section
9	201(ff) of the Federal Food, Drug, and Cosmetic
10	Act (21 U.S.C. 321(f)).

11 SEC. 8. PENALTY.

A health care practitioner who knowingly violates any
provision of this Act shall not be covered by the protections under this Act and shall be subject to all other applicable laws and regulations.

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