111TH CONGRESS 2D SESSION

H.R. 1032

AN ACT

To amend the Public Health Service Act to improve the prevention, diagnosis, and treatment of heart disease, stroke, and other cardiovascular diseases in women.

- 1 Be it enacted by the Senate and House of Representa-
- ${\it 2\ tives\ of\ the\ United\ States\ of\ America\ in\ Congress\ assembled},$

1 SECTION 1. SHORT TITLE.

- This Act may be cited as the "Heart Disease Edu-
- 3 cation, Analysis Research, and Treatment for Women
- 4 Act" or the "HEART for Women Act".
- 5 SEC. 2. REPORT BY GOVERNMENT ACCOUNTABILITY OF-
- 6 FICE.
- 7 (a) In General.—The Comptroller General of the
- 8 United States shall conduct a study investigating the ex-
- 9 tent to which sponsors of clinical studies of investigational
- 10 drugs, biologics, and devices and sponsors of applications
- 11 for approval or licensure of new drugs, biologics, and de-
- 12 vices comply with Food and Drug Administration require-
- 13 ments and follow guidance for presentation of clinical
- 14 study safety and effectiveness data by sex, age, and racial
- 15 subgroups.
- 16 (b) Report by GAO.—
- 17 (1) Submission.—Not later than 12 months
- after the date of the enactment of this Act, the
- 19 Comptroller General shall complete the study under
- subsection (a) and submit to the Committee on En-
- 21 ergy and Commerce of the House of Representatives
- and the Committee on Health, Education, Labor,
- and Pensions of the Senate a report on the results
- of such study.
- 25 (2) Contents.—The report required by para-
- 26 graph (1) shall include each of the following:

1	(A) A description of the extent to which						
2	the Food and Drug Administration assists						
3	sponsors in complying with the requirements						
4	and following the guidance referred to in sub-						
5	section (a).						
6	(B) A description of the effectiveness of						
7	the Food and Drug Administration's enforce-						
8	ment of compliance with such requirements.						
9	(C) An analysis of the extent to which fe-						
10	males, racial and ethnic minorities, and adults						
11	of all ages are adequately represented in Food						
12	and Drug Administration-approved clinical						
13	studies (at all phases) so that product safety						
14	and effectiveness data can be evaluated by gen-						
15	der, age, and racial subgroup.						
16	(D) An analysis of the extent to which a						
17	summary of product safety and effectiveness						
18	data disaggregated by sex, age, and racial sub-						
19	group is readily available to the public in a						
20	timely manner by means of the product label or						
21	the Food and Drug Administration's Website.						
22	(E) Appropriate recommendations for—						
23	(i) modifications to the requirements						
24	and guidance referred to in subsection (a);						
25	or						

1	(ii) oversight by the Food and Drug
2	Administration of such requirements.
3	(c) REPORT BY HHS.—Not later than 6 months
4	after the submission by the Comptroller General of the
5	report required under subsection (b), the Secretary of
6	Health and Human Services shall submit to the Com-
7	mittee on Energy and Commerce of the House of Rep-
8	resentatives and the Committee on Health, Education,
9	Labor, and Pensions of the Senate a response to that re-
10	port, including a corrective action plan as needed to re-
11	spond to the recommendations in that report.
12	(d) DEFINITIONS.—In this section:
13	(1) The term "biologic" has the meaning given
14	to the term "biological product" in section 351(i) of
15	the Public Health Service Act (42 U.S.C. 262(i)).
16	(2) The term "device" has the meaning given to
17	such term in section 201(h) of the Federal Food,
18	Drug, and Cosmetic Act (21 U.S.C. 321(h)).
19	(3) The term "drug" has the meaning given to
20	such term in section 201(g) of the Federal Food,
21	Drug, and Cosmetic Act (21 U.S.C. 321(g)).

1	SEC. 3. REPORTING ON QUALITY OF AND ACCESS TO CARE
2	FOR WOMEN WITH CARDIOVASCULAR DIS-
3	EASES.
4	Part P of title III of the Public Health Service Act
5	(42 U.S.C. 280g et seq.) is amended by adding at the end
6	the following:
7	"SEC. 399V-5. REPORTING ON QUALITY OF AND ACCESS TO
8	CARE FOR WOMEN WITH CARDIOVASCULAR
9	DISEASES.
10	"Not later than September 30, 2013, and annually
11	thereafter, the Secretary of Health and Human Services
12	shall prepare and submit to the Congress a report on the
13	quality of and access to care for women with heart disease,
14	stroke, and other cardiovascular diseases. The report shall
15	contain recommendations for eliminating disparities in,
16	and improving the treatment of, heart disease, stroke, and
17	other cardiovascular diseases in women.".
18	SEC. 4. EXTENSION OF WISEWOMAN PROGRAM.
19	Section 1509 of the Public Health Service Act (42
20	U.S.C. 300n-4a) is amended—
21	(1) in subsection (a)—
22	(A) by striking the heading and inserting
23	"In General.—"; and
24	(B) in the matter preceding paragraph (1),
25	by striking "may make grants" and all that fol-
26	lows through "purpose" and inserting the fol-

1	lowing: "may make grants to such States for
2	the purpose"; and
3	(2) in subsection (d)(1), by striking "there are
4	authorized" and all that follows through the period
5	and inserting "there are authorized to be appro-
6	priated \$23,000,000 for fiscal year 2012,
7	\$25,300,000 for fiscal year 2013, $$27,800,000$ for
8	fiscal year 2014, $$30,800,000$ for fiscal year 2015,
9	and \$34,000,000 for fiscal year 2016.".
	Passed the House of Representatives September 30

Attest:

(legislative day September 29), 2010.

Clerk.

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