

CERTIFICATION OF ENROLLMENT
SECOND SUBSTITUTE HOUSE BILL 1745

68th Legislature
2023 Regular Session

Passed by the House April 19, 2023
Yeas 98 Nays 0

**Speaker of the House of
Representatives**

Passed by the Senate April 12, 2023
Yeas 49 Nays 0

President of the Senate

Approved

Governor of the State of Washington

CERTIFICATE

I, Bernard Dean, Chief Clerk of the House of Representatives of the State of Washington, do hereby certify that the attached is **SECOND SUBSTITUTE HOUSE BILL 1745** as passed by the House of Representatives and the Senate on the dates hereon set forth.

Chief Clerk

FILED

**Secretary of State
State of Washington**

SECOND SUBSTITUTE HOUSE BILL 1745

AS AMENDED BY THE SENATE

Passed Legislature - 2023 Regular Session

State of Washington 68th Legislature 2023 Regular Session

By House Appropriations (originally sponsored by Representatives Thai, Duerr, Doglio, Ormsby, and Macri)

READ FIRST TIME 02/24/23.

1 AN ACT Relating to improving diversity in clinical trials;
2 amending RCW 43.348.040; adding a new section to chapter 43.348 RCW;
3 adding a new section to chapter 28B.20 RCW; adding a new section to
4 chapter 28B.30 RCW; adding a new chapter to Title 69 RCW; creating a
5 new section; and providing an expiration date.

6 BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF WASHINGTON:

7 NEW SECTION. **Sec. 1.** (1) The legislature finds that controlled
8 clinical trials provide a critical base of evidence for evaluating
9 whether a medical product is safe and effective before the product is
10 approved for marketing. The United States food and drug
11 administration has evaluated demographic profiles of people
12 participating in clinical trials for approved drugs and found that
13 some groups, especially ethnic and racial groups, are not always well
14 represented in clinical trials. Diversity in clinical trials is
15 necessary to effectively determine how race, gender, and age impact
16 how a person metabolizes a drug. Communities of color have been
17 working diligently to establish a foundation of trust with government
18 and clinical research with the goal of engaging more trial
19 participants who are members of underrepresented demographic groups.
20 Joining clinical trials is a difficult and complex process and the
21 lack of trust and awareness of clinical trials and research, in

1 addition to burdens related to transportation, geography, and access,
2 limit trial participants. The lack of diversity in clinical trials
3 compounds access to treatment disparities and limits our
4 understanding of the impacts of studied interventions and conditions
5 across the population.

6 (2) Therefore, it is the policy of the state to:

7 (a) Improve the completeness and quality of data concerning
8 diverse demographic groups that is collected, reported, and analyzed
9 for the purposes of clinical trials of drugs and medical devices;

10 (b) Identify barriers to participation in clinical trials by
11 persons who are members of demographic groups that are
12 underrepresented in such trials and employ strategies recognized by
13 the United States food and drug administration to encourage greater
14 participation in clinical trials by such persons;

15 (c) Make data concerning demographic groups that is collected,
16 reported, and analyzed for the purposes of clinical trials more
17 available and transparent; and

18 (d) Require certain entities conducting clinical trials to offer
19 trial participants information in a language other than English and
20 provide culturally specific recruitment materials alongside general
21 enrollment materials.

22 NEW SECTION. **Sec. 2.** The definitions in this section apply
23 throughout this chapter unless the context clearly requires
24 otherwise.

25 (1) "Washington state review board" or "review board" means the
26 Washington state institutional review board, established pursuant to
27 45 C.F.R. Part 46, which is the designated institutional review board
28 for the department of social and health services, the department of
29 health, the department of labor and industries, and other state
30 agencies.

31 (2) "Underrepresented community" or "underrepresented demographic
32 group" means a community or demographic group that is more likely to
33 be historically marginalized and less likely to be included in
34 research and clinical trials represented by race, sex, sexual
35 orientation, socioeconomic status, age, and geographic location.

36 NEW SECTION. **Sec. 3.** The Washington state review board shall
37 establish a diversity in clinical trials program to encourage
38 participation in clinical trials of drugs and medical devices by

1 persons who are members of demographic groups that are
2 underrepresented in clinical trials. In developing this program, the
3 review board shall compile and share information and resources in an
4 accessible fashion to assist entities in Washington state that
5 conduct clinical trials of drugs and medical devices to increase
6 participation by persons who are members of demographic groups that
7 are underrepresented in clinical trials including, but not limited
8 to:

9 (1) Information concerning methods for identifying and recruiting
10 persons who are members of underrepresented demographic groups to
11 participate in clinical trials;

12 (2) Links or copies of outside resources related to increasing
13 participation by members of underrepresented demographic groups in
14 clinical trials provided by community organizations or other
15 interested agencies or parties;

16 (3) Contact information for community organizations or other
17 appropriate entities which may be able to provide assistance with
18 efforts to increase participation by underrepresented demographic
19 groups in clinical trials; and

20 (4) Links to websites maintained by medical facilities, health
21 authorities, and other local governmental entities, nonprofit
22 organizations, and scientific investigators and institutions that are
23 performing research relating to drugs or medical devices in this
24 state.

25 NEW SECTION. **Sec. 4.** Any state entity or hospital that receives
26 funding from the national institutes of health to conduct clinical
27 trials of drugs or medical devices shall:

28 (1) Adopt a policy concerning the identification and recruitment
29 of persons who are members of underrepresented demographic groups to
30 participate in clinical trials. This policy must include requirements
31 that investigators who are conducting clinical trials collaborate
32 with community-based organizations and use methods recognized by the
33 United States food and drug administration to identify and recruit
34 such persons to participate in those clinical trials;

35 (2) Provide information to trial participants in languages other
36 than English;

37 (3) Provide translation services or bilingual staff for trial
38 screening;

1 (4) Provide culturally specific recruitment materials alongside
2 general enrollment materials; and

3 (5) Provide electronic consent when not prohibited by the
4 granting entity or federal regulations.

5 **Sec. 5.** RCW 43.348.040 and 2018 c 4 s 4 are each amended to read
6 as follows:

7 (1) The Andy Hill cancer research endowment program is created.
8 The purpose of the program is to make grants to public and private
9 entities, including commercial entities, to fund or reimburse the
10 entities pursuant to agreement for the promotion of cancer research
11 to be conducted in the state. The endowment is to oversee and guide
12 the program, including the solicitation, selection, and award of
13 grants.

14 (2) The board must develop a plan for the allocation of projected
15 amounts in the fund, which it must update annually, following at
16 least one annual public hearing. The plan must provide for
17 appropriate funding continuity and take into account the projected
18 speed at which revenues will be available and amounts that can be
19 spent during the plan period.

20 (3) The endowment must solicit requests for grant funding and
21 evaluate the requests by reference to factors such as: (a) The
22 quality of the proposed research or program; (b) its potential to
23 improve health outcomes of persons with cancer, with particular
24 attention to the likelihood that it will also lower health care
25 costs, substitute for a more costly diagnostic or treatment modality,
26 or offer a breakthrough treatment for a particular cancer or cancer-
27 related condition or disease; (c) its potential for leveraging
28 additional funding; (d) its potential to provide additional health
29 care benefits or benefit other human diseases or conditions; (e) its
30 potential to stimulate life science, health care, and biomedical
31 employment in the state; (f) the geographic diversity of the grantees
32 within Washington; (g) evidence of potential royalty, sales, or
33 licensing revenue, or other commercialization-related revenue and
34 contractual means to recapture such income for purposes of this
35 chapter; ~~((and))~~ (h) evidence of public and private collaboration;
36 (i) the ability to offer trial participants information in a language
37 other than English; (j) the ability to provide culturally specific
38 recruitment materials alongside general enrollment materials; (k) the
39 ability to provide electronic consent when not prohibited by other

1 granting entities or federal regulations; and (l) other evidence of
2 outreach and engagement to increase participation of underrepresented
3 communities in clinical trials of drugs and medical devices.

4 (4) The endowment may not award a grant for a proposal that was
5 not recommended by an independent expert scientific review and
6 advisory committee under RCW 43.348.050.

7 (5) The endowment must issue an annual report to the public that
8 sets forth its activities with respect to the fund, including grants
9 awarded, grant-funded work in progress, research accomplishments,
10 prevention, and care activities, and future program directions with
11 respect to cancer research, prevention, and care. Each annual report
12 regarding activities of the program and fund must include, but not be
13 limited to, the following: The number and dollar amounts of grants;
14 the grantees for the prior year; the endowment's administrative
15 expenses; an assessment of the availability of funding for cancer
16 research, prevention, and care from sources other than the endowment;
17 a summary of research, prevention, and care-related findings,
18 including promising new areas for investment; and a report on the
19 benefits to Washington of its programs to date.

20 (6) The endowment's first annual report must include a proposed
21 operating plan for the design, implementation, and administration of
22 an endowment program supporting the purposes of the endowment and
23 program.

24 (7) The endowment must adopt policies to ensure that all
25 potential conflicts have been disclosed and that all conflicts have
26 been eliminated or mitigated.

27 (8) The endowment must establish standards to ensure that
28 recipients of grants for cancer research, prevention, or care
29 purchase goods and services from Washington suppliers to the extent
30 reasonably possible.

31 NEW SECTION. **Sec. 6.** (1) The department of health, in
32 consultation with the University of Washington, Washington State
33 University, the Andy Hill cancer research endowment, Washington
34 community health boards and initiatives, community-based
35 organizations, and other relevant research organizations, shall
36 analyze and provide recommendations on the following:

37 (a) What demographic groups and populations are currently
38 represented and underrepresented in clinical trials in Washington,
39 including geographic representation;

1 (b) Barriers for persons who are members of underrepresented
2 demographic groups to participate in clinical trials in Washington,
3 including barriers related to transportation; and

4 (c) Approaches for how clinical trials can successfully partner
5 with community-based organizations and others to provide outreach to
6 underrepresented communities.

7 (2) By December 1, 2023, the department of health shall report to
8 the legislature the results of the analysis and any recommendations
9 to increase diversity and reduce barriers for participants in
10 clinical trials.

11 (3) For purposes of this section, "underrepresented community" or
12 "underrepresented demographic group" means a community or demographic
13 group that is more likely to be historically marginalized and less
14 likely to be included in research and clinical trials represented by
15 race, sex, sexual orientation, socioeconomic status, age, and
16 geographic location.

17 (4) This section expires December 31, 2023.

18 NEW SECTION. **Sec. 7.** A new section is added to chapter 43.348
19 RCW to read as follows:

20 (1) Beginning January 1, 2024, the University of Washington and
21 Washington State University may partner with the Andy Hill cancer
22 research endowment, the department of health, community-based
23 organizations, and other entities to increase the participation of
24 persons who are members of underrepresented demographic groups in
25 clinical trials for drugs or medical devices. If an investigator at
26 the University of Washington or Washington State University is
27 conducting or planning to conduct a clinical trial on a drug or
28 medical device and the University determines that the trial would
29 benefit from specific community outreach and engagement to increase
30 participation of an underrepresented community in the clinical trial,
31 the University of Washington or Washington State University may:

32 (a) Request the assistance of the department of health and the
33 Andy Hill cancer research endowment to create an outreach plan and
34 coordinate with community-based organizations to provide outreach and
35 engagement; and

36 (b) Provide the Andy Hill cancer research endowment and the
37 department of health with the following information:

1 (i) A summary of the clinical trial, including a description of
2 the drug or medical device and any condition or disease that the
3 clinical trial is addressing or targeting;

4 (ii) Any information on health disparities related to the
5 condition, disease, or related drugs or medical devices, including
6 any demographic groups that may be disproportionately impacted; and

7 (iii) Any other information that may assist the Andy Hill cancer
8 research endowment, department of health, and community-based
9 organizations in providing outreach and engagement to specific
10 demographic groups or communities.

11 (2) The requesting university, the Andy Hill cancer research
12 endowment, and the department of health, in collaboration with
13 community-based organizations and other appropriate entities, shall
14 develop a specific community outreach and engagement plan to increase
15 participation of an underrepresented demographic group or community
16 in the clinical trial.

17 (3) Subject to the availability of amounts appropriated for this
18 specific purpose, the Andy Hill cancer research endowment may
19 administer grants to Washington state community-based organizations
20 to implement the outreach plan and to provide meaningful and real-
21 time community engagement with any demographic groups or communities
22 identified in subsection (1) of this section with the goal of
23 increasing the demographic group's or community's participation in
24 the clinical trial. The community engagement should utilize any
25 recommendations provided by the department of health's report
26 required under section 6 of this act.

27 NEW SECTION. **Sec. 8.** A new section is added to chapter 28B.20
28 RCW to read as follows:

29 If at any time the University of Washington receives funding from
30 the national institutes of health to conduct clinical trials of drugs
31 or medical devices, the University of Washington shall adopt a policy
32 concerning the identification and recruitment of persons who are
33 members of underrepresented demographic groups to participate in
34 clinical trials of drugs and medical devices. This policy must
35 include requirements to:

36 (1) Adopt a policy concerning the identification and recruitment
37 of persons who are members of underrepresented demographic groups to
38 participate in clinical trials. This policy must include requirements
39 that investigators who are conducting clinical trials collaborate

1 with community-based organizations and use methods recognized by the
2 United States food and drug administration to identify and recruit
3 such persons to participate in those clinical trials;

4 (2) Provide information to trial participants in languages other
5 than English;

6 (3) Provide translation services or bilingual staff for trial
7 screening;

8 (4) Provide culturally specific recruitment materials alongside
9 general enrollment materials; and

10 (5) Provide electronic consent when not prohibited by the
11 granting entity or federal regulations.

12 NEW SECTION. **Sec. 9.** A new section is added to chapter 28B.30
13 RCW to read as follows:

14 If at any time Washington State University receives funding from
15 the national institutes of health to conduct clinical trials of drugs
16 or medical devices, Washington State University shall adopt a policy
17 concerning the identification and recruitment of persons who are
18 members of underrepresented demographic groups to participate in
19 clinical trials of drugs and medical devices. This policy must
20 include requirements to:

21 (1) Adopt a policy concerning the identification and recruitment
22 of persons who are members of underrepresented demographic groups to
23 participate in clinical trials. This policy must include requirements
24 that investigators who are conducting clinical trials collaborate
25 with community-based organizations and use methods recognized by the
26 United States food and drug administration to identify and recruit
27 such persons to participate in those clinical trials;

28 (2) Provide information to trial participants in languages other
29 than English;

30 (3) Provide translation services or bilingual staff for trial
31 screening;

32 (4) Provide culturally specific recruitment materials alongside
33 general enrollment materials; and

34 (5) Provide electronic consent when not prohibited by the
35 granting entity or federal regulations.

1 NEW SECTION. **Sec. 10.** Sections 1 through 4 of this act
2 constitute a new chapter in Title 69 RCW.

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