119TH CONGRESS	$\mathbf{C}$	
1st Session		
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To amend section 495 of the Public Health Service Act to require inspections of foreign laboratories conducting biomedical and behavioral research to ensure compliance with applicable animal welfare requirements, and for other purposes

## IN THE SENATE OF THE UNITED STATES

Mr. Schmitt (for himself, Mr. Merkley, Mr. Ricketts, Mr. Fetterman, Mr. Scott of Florida, Mr. Peters, and Ms. Ernst) introduced the following bill; which was read twice and referred to the Committee on

## A BILL

To amend section 495 of the Public Health Service Act to require inspections of foreign laboratories conducting biomedical and behavioral research to ensure compliance with applicable animal welfare requirements, 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 TITLES.
- 4 This Act may be cited as the "Worldwide Animal
- 5 Testing Compliance and Harmonization Act of 2025" or
- 6 the "WATCH Act".

1	SEC. 2. FOREIGN LABORATORY INSPECTIONS AND CERTIFI-
2	CATION.
3	(a) In General.—Section 495 of the Public Health
4	Service Act (42 U.S.C. 289d) is amended by adding at
5	the end the following:
6	"(f) Inspection and Certification of Foreign
7	Laboratories.—
8	"(1) In general.—As a condition of eligibility
9	to perform research involving animals under a grant,
10	contract, or cooperative agreement administered by
11	the National Institutes of Health or any national re-
12	search institute, a laboratory located outside the
13	United States that receives Federal funds shall be
14	subject to quarterly inspections to evaluate compli-
15	ance with the requirements under this title.
16	"(2) Inspection and Certification Re-
17	QUIREMENTS.—
18	"(A) QUARTERLY INSPECTION PROCESS.—
19	The Secretary, in consultation with appropriate
20	foreign regulatory authorities and international
21	organizations, shall establish and implement a
22	process for conducting quarterly inspections of
23	foreign laboratories that have received an Ani-
24	mal Welfare Assurance (as defined in section

9.2 of title 42, Code of Federal Regulations) to

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1	ensure their continued compliance with the re-
2	quirements under this title.
3	"(B) Assurances.—The inspection proc-
4	ess established by the Secretary pursuant to
5	subparagraph (A) shall evaluate the compliance
6	of foreign laboratories with the requirements
7	under subsection (c)(1), including—
8	"(i) the establishment and operation
9	of animal care committees;
10	"(ii) the review and evaluation of ani-
11	mal care and treatment; and
12	"(iii) proper record-keeping and re-
13	porting procedures.
14	"(3) Certification of compliance and pub-
15	LIC ACCESS.—
16	"(A) Issuance.—Following each quarterly
17	inspection required under paragraph (2)(A), the
18	inspecting authority shall issue a certification of
19	compliance to the laboratories determined to be
20	in compliance with the requirements under
21	paragraph (2)(B).
22	"(B) Public Access.—Copies of the cer-
23	tificates of compliance issued pursuant to sub-
24	paragraph (A) shall be maintained by the Office
25	of Laboratory Animal Welfare and shall remain

1	publicly accessible with other information about
2	currently issued Animal Welfare Assurances.
3	"(C) Corrective action.—Laboratories
4	that fail to comply with the requirements under
5	paragraph (2)(B) shall be given a reasonable
6	opportunity to take corrective action.
7	"(4) Suspension or revocation of grant
8	OR CONTRACT FOR NON-COMPLIANT FOREIGN LAB
9	ORATORIES.—If the Secretary determines that a for-
10	eign facility is not in compliance with the require-
11	ments under subsection $(c)(1)$ and does not take ap-
12	propriate corrective action after given a reasonable
13	opportunity to do so, the Secretary shall suspend or
14	revoke the applicable grant, contract, or cooperative
15	agreement involving research on animals under such
16	conditions as the Director of NIH determines appro-
17	priate, in accordance with subsection (d).
18	"(5) Designation of Inspecting Author
19	ITY.—The Secretary, in consultation with the Direc-
20	tor of NIH, shall designate an appropriate authority
21	to conduct the quarterly inspections required under
22	paragraph (2)(A) and issue certifications of compli-
23	ance in accordance with paragraph (3).
24	"(6) Coordination with foreign authoris
25	TIES.—The Secretary and the Director of NIH shal

- 1 coordinate with appropriate foreign regulatory au-
- 2 thorities and enter into agreements with foreign gov-
- 3 ernments, as needed, to facilitate the implementation
- 4 and enforcement of this subsection, while respecting
- 5 the sovereignty and laws of foreign nations.".
- 6 (b) Effective Date.—The amendment made by
- 7 subsection (a) shall take effect on the date that is 180
- 8 days after the date of the enactment of this Act.