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2D SESSION

# H. R. 8333

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IN THE SENATE OF THE UNITED STATES

SEPTEMBER 10, 2024

Received; read twice and referred to the Committee on Homeland Security and  
Governmental Affairs

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## AN ACT

To prohibit contracting with certain biotechnology providers,  
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,*

## **1 SECTION 1. SHORT TITLE.**

2 This Act may be cited as the “BIOSECURE Act”.

### **3 SEC. 2. PROHIBITION ON CONTRACTING WITH CERTAIN**

## 4 BIOTECHNOLOGY PROVIDERS.

5 (a) IN GENERAL.—The head of an executive agency

## 6 may not—

7                   (1) procure or obtain any biotechnology equip-  
8       ment or service produced or provided by a bio-  
9       technology company of concern; or

(2) enter into a contract or extend or renew a  
contract with any entity that—

(B) enters into any contract the performance of which such entity knows or has reason to believe will require, in performance of the contract with the executive agency, the use of biotechnology equipment or services produced or provided by a biotechnology company of concern and acquired after the applicable effective date in subsection (c).

1       (b) PROHIBITION ON LOAN AND GRANT FUNDS.—

2   The head of an executive agency may not obligate or ex-  
3  pend loan or grant funds to, and a loan or grant recipient  
4  may not use loan or grant funds to—

5           (1) procure, obtain, or use any biotechnology  
6  equipment or services produced or provided by a bio-  
7  technology company of concern; or

8           (2) enter into a contract or extend or renew a  
9  contract with an entity described in subsection  
10 (a)(2).

11       (c) EFFECTIVE DATES.—

12           (1) CERTAIN ENTITIES.—With respect to the  
13  biotechnology companies of concern covered by sub-  
14  section (f)(2)(A), the prohibitions under subsections  
15  (a) and (b) shall take effect 60 days after the  
16  issuance of the regulation in subsection (h).

17           (2) OTHER ENTITIES.—With respect to the bio-  
18  technology companies of concern covered by sub-  
19  section (f)(2)(B), the prohibitions under subsections  
20  (a) and (b) shall take effect 180 days after the  
21  issuance of the regulation in subsection (h).

22           (3) RULES OF CONSTRUCTION.—

23           (A) CERTAIN ENTITIES.—Prior to January  
24  1, 2032, with respect to biotechnology compa-  
25  nies of concern covered by subsections

1                         (f)(2)(A), subsections (a)(2) and (b)(2) shall  
2                         not apply to biotechnology equipment or serv-  
3                         ices produced or provided under a contract or  
4                         agreement, including previously negotiated con-  
5                         tract options, entered into before the effective  
6                         date under paragraph (1).

7                         (B) OTHER ENTITIES.—Prior to the date  
8                         that is five years after the issuance of the regu-  
9                         lation in subsection (h) that identifies a bio-  
10                         technology company of concern covered by sub-  
11                         sections (f)(2)(B), subsections (a)(2) and (b)(2)  
12                         shall not apply to biotechnology equipment or  
13                         services produced or provided under a contract  
14                         or agreement, including previously negotiated  
15                         contract options, entered into before the effec-  
16                         tive date under paragraph (2).

17                         (C) SAFE HARBOR.—The term “bio-  
18                         technology equipment or services produced or  
19                         provided by a biotechnology company of con-  
20                         cern” shall not be construed to refer to any bio-  
21                         technology equipment or services that were for-  
22                         merly, but are no longer, produced or provided  
23                         by biotechnology companies of concern.

24                         (d) WAIVER AUTHORITIES.—

25                         (1) SPECIFIC BIOTECHNOLOGY EXCEPTION.—

(A) WAIVER.—The head of the applicable executive agency may waive the prohibition under subsections (a) and (b) on a case-by-case basis—

(i) with the approval of the Director of the Office of Management and Budget, in coordination with the Secretary of Defense; and

(ii) if such head submits a notification and justification to the appropriate congressional committees not later than 30 days after granting such waiver.

(B) DURATION.—

(i) IN GENERAL.—Except as provided in clause (ii), a waiver granted under subparagraph (A) shall last for a period of not more than 365 days.

(ii) EXTENSION.—The head of the applicable executive agency, with the approval of the Director of the Office of Management and Budget, and in coordination with the Secretary of Defense, may extend a waiver granted under subparagraph (A) one time, for a period up to 180 days after the date on which the waiver

1           would otherwise expire, if such an exten-  
2           sion is in the national security interests of  
3           the United States and if such head sub-  
4           mits a notification and justification to the  
5           appropriate congressional committees not  
6           later than 10 days after granting such  
7           waiver extension.

8           (2) OVERSEAS HEALTH CARE SERVICES.—The  
9           head of an executive agency may waive the prohibi-  
10          tions under subsections (a) and (b) with respect to  
11          a contract, subcontract, or transaction for the acqui-  
12          sition or provision of health care services overseas on  
13          a case-by-case basis—

14           (A) if the head of such executive agency  
15          determines that the waiver is—

16               (i) necessary to support the mission or  
17          activities of the employees of such execu-  
18          tive agency described in subsection  
19          (e)(2)(A); and

20               (ii) in the interest of the United  
21          States;

22           (B) with the approval of the Director of  
23          the Office of Management and Budget, in con-  
24          sultation with the Secretary of Defense; and

5       (e) EXCEPTIONS.—The prohibitions under sub-  
6 sections (a) and (b) shall not apply to—

(B) employees of contractors or sub-contractors of the United States—

21 (i) who are performing under a con-  
22 tract that directly supports the missions or  
23 activities of individuals described in sub-  
24 paragraph (A); and

7       (f) EVALUATION OF CERTAIN BIOTECHNOLOGY EN-  
8 TITIES.—

(A) BGI, MGI, Complete Genomics, WuXi AppTec, and WuXi Biologics;

(B) any entity that is determined by the process established in paragraph (1) to meet the following criteria—

(i) is subject to the administrative governance structure, direction, control, or operates on behalf of the government of a foreign adversary;

(ii) is to any extent involved in the manufacturing, distribution, provision, or procurement of a biotechnology equipment or service; and

(iii) poses a risk to the national security of the United States based on—

(I) engaging in joint research with, being supported by, or being affiliated with a foreign adversary's military, internal security forces, or intelligence agencies;

(II) providing multiomic data obtained via biotechnology equipment or services to the government of a foreign adversary; or

(III) obtaining human multiomic data via the biotechnology equipment

1                   or services without express and in-  
2                   formed consent; and

3                   (C) any subsidiary, parent, affiliate, or  
4                   successor of entities listed in subparagraphs (A)  
5                   and (B), provided they meet the criteria in sub-  
6                   paragraph (B)(i).

7                   (3) GUIDANCE.—Not later than 120 days after  
8                   the date of the enactment of this Act for the bio-  
9                   technology companies of concern named in para-  
10                  graph (2)(A), and not later than 180 days after the  
11                  development of the list pursuant to paragraph (1)  
12                  and any update to the list pursuant to paragraph  
13                  (4), the Director of the Office of Management and  
14                  Budget, in coordination with the Secretary of De-  
15                  fense, the Attorney General, the Secretary of Health  
16                  and Human Services, the Secretary of Commerce,  
17                  the Director of National Intelligence, the Secretary  
18                  of Homeland Security, the Secretary of State, and  
19                  the National Cyber Director, shall establish guidance  
20                  as necessary to implement the requirements of this  
21                  section.

22                  (4) UPDATES.—The Director of the Office of  
23                  Management and Budget, in coordination with or  
24                  based on a recommendation provided by the Sec-  
25                  retary of Defense, the Attorney General, the Sec-

1           retary of Health and Human Services, the Secretary  
2           of Commerce, the Director of National Intelligence,  
3           the Secretary of Homeland Security, the Secretary  
4           of State, and the National Cyber Director, shall pe-  
5           riodically, though not less than annually, review and,  
6           as appropriate, modify the list of biotechnology com-  
7           panies of concern, and notify the appropriate con-  
8           gressional committees of any such modifications.

9                         (5) NOTICE OF A DESIGNATION AND REVIEW.—

10                         (A) IN GENERAL.—A notice of a designa-  
11                         tion as a biotechnology company of concern  
12                         under paragraph (2)(B) shall be issued to any  
13                         biotechnology company of concern named in the  
14                         designation—

15                                 (i) advising that a designation has  
16                         been made;

17                                 (ii) identifying the criteria relied upon  
18                         under such subparagraph and, to the ex-  
19                         tent consistent with national security and  
20                         law enforcement interests, the information  
21                         that formed the basis for the designation;

22                                 (iii) advising that, within 90 days  
23                         after receipt of notice, the biotechnology  
24                         company of concern may submit informa-

1                      tion and argument in opposition to the  
2                      designation;

3 (iv) describing the procedures gov-  
4 erning the review and possible issuance of  
5 a designation pursuant to paragraph (1);  
6 and

12 (B) CONGRESSIONAL NOTIFICATION RE-  
13 QUIREMENTS.—

ignation pursuant to subparagraph (A)(iii),  
the Director of the Office of Management  
and Budget shall submit such information  
to the Committee on Homeland Security  
and Governmental Affairs of the Senate  
and the Committee on Oversight and Ac-  
countability of the House of Representa-  
tives.

21 (g) EVALUATION OF NATIONAL SECURITY RISKS  
22 POSED BY FOREIGN ADVERSARY ACQUISITION OF AMER-  
23 ICAN MULTIMIC DATA.—

1       tional Intelligence, in consultation with the Secretary  
2       of Defense, the Attorney General of the United  
3       States, the Secretary of Health and Human Serv-  
4       ices, the Secretary of Commerce, the Secretary of  
5       Homeland Security, the Secretary of State, and the  
6       National Cyber Director, shall complete an assess-  
7       ment of risks to national security posed by human  
8       multiomic data from United States citizens that is  
9       collected or stored by a foreign adversary from the  
10      provision of biotechnology equipment or services.

11                   (2) REPORT REQUIREMENT.—Not later than 30  
12      days after the completion of the assessment devel-  
13      oped under paragraph (1), the Director of National  
14      Intelligence shall submit a report with such assess-  
15      ment to the appropriate congressional committees.

16                   (3) FORM.—The report required under para-  
17      graph (2) shall be in unclassified form accompanied  
18      by a classified annex.

19                   (h) REGULATIONS.—Not later than one year after  
20      the date of establishment of guidance required under sub-  
21      section (f)(3), and as necessary for subsequent updates,  
22      the Federal Acquisition Regulatory Council shall revise  
23      the Federal Acquisition Regulation as necessary to imple-  
24      ment the requirements of this section.

1           (i) REPORTING ON INTELLIGENCE ON NEFARIOUS  
2 ACTIVITIES OF BIOTECHNOLOGY COMPANIES WITH  
3 HUMAN MULTIOMIC DATA.—Not later than 180 days  
4 after the date of the enactment of this Act, and annually  
5 thereafter, the Director of National Intelligence, in con-  
6 sultation with the heads of executive agencies, shall submit  
7 to the appropriate congressional committees a report on  
8 any intelligence in possession of such agencies related to  
9 nefarious activities conducted by biotechnology companies  
10 with human multiomic data. The report shall include in-  
11 formation pertaining to potential threats to national secu-  
12 rity or public safety from the selling, reselling, licensing,  
13 trading, transferring, sharing, or otherwise providing or  
14 making available to any foreign country of any forms of  
15 multiomic data of a United States citizen.

16           (j) NO ADDITIONAL FUNDS.—No additional funds  
17 are authorized to be appropriated for the purpose of car-  
18 rying out this section.

19           (k) DEFINITIONS.—In this section:

20               (1) APPROPRIATE CONGRESSIONAL COMMIT-  
21 TEES.—The term “appropriate congressional com-  
22 mittees” means—  
23                       (A) the Committee on Armed Services, the  
24 Select Committee on Intelligence, and the Com-

1           mittee on Homeland Security and Govern-  
2           mental Affairs of the Senate; and

3                 (B) the Committee on Armed Services, the  
4                 Permanent Select Committee on Intelligence,  
5                 the Committee on Foreign Affairs, the Com-  
6                 mittee on Oversight and Accountability, the  
7                 Committee on Energy and Commerce, and the  
8                 Select Committee on Strategic Competition be-  
9                 tween the United States and the Chinese Com-  
10                munist Party of the House of Representatives.

11                 (2) BIOTECHNOLOGY EQUIPMENT OR SERV-  
12                 ICE.—The term “biotechnology equipment or serv-  
13                 ice” means—

14                         (A) equipment, including genetic sequenc-  
15                 ers, combined mass spectrometry technologies,  
16                 polymerase chain reaction machines, or any  
17                 other instrument, apparatus, machine, or de-  
18                 vice, including components and accessories  
19                 thereof, that is designed for use in the research,  
20                 development, production, or analysis of biologi-  
21                 cal materials as well as any software, firmware,  
22                 or other digital components that are specifically  
23                 designed for use in, and necessary for the oper-  
24                 ation of, such equipment;

(ii) disease detection, genealogical information, and related services; and

(3) CONTRACT.—Except as the term is used under subsection (b)(2) and subsection (c)(3), the

1       term “contract” means any contract subject to the  
2       Federal Acquisition Regulation issued under section  
3       1303(a)(1) of title 41, United States Code.

4                 (4) CONTROL.—The term “control” has the  
5       meaning given to that term in section 800.208 of  
6       title 31, Code of Federal Regulations, or any suc-  
7       cessor regulations.

8                 (5) EXECUTIVE AGENCY.—The term “executive  
9       agency” has the meaning given the term “Executive  
10      agency” in section 105 of title 5, United States  
11      Code.

12                 (6) FOREIGN ADVERSARY.—The term “foreign  
13       adversary” has the meaning given the term “covered  
14       nation” in section 4872(d) of title 10, United States  
15       Code.

16                 (7) MULTIOMIC.—The term “multiomic” means  
17       data types that include genomics, epigenomics,  
18       transcriptomics, proteomics, and metabolomics.

19                 (8) OVERSEAS.—The term “overseas” means  
20       any area outside of the United States, the Common-

1       wealth of Puerto Rico, or a territory or possession  
2       of the United States.

Passed the House of Representatives September 9,  
2024.

Attest:                   KEVIN F. MCCUMBER,  
*Clerk.*