

119TH CONGRESS  
2D SESSION

# H. R. 9102

To amend title VIII of the Defense Production Act of 1950 to alter the definitions of “prohibited technology” and “notifiable technology”, and for other purposes.

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## IN THE HOUSE OF REPRESENTATIVES

JUNE 2, 2026

Mr. MOOLENAAR (for himself and Mrs. DINGELL) introduced the following bill; which was referred to the Committee on Financial Services

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## A BILL

To amend title VIII of the Defense Production Act of 1950 to alter the definitions of “prohibited technology” and “notifiable technology”, 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 at the “Biotech Investment  
5       National Security Act of 2026” or the “BINS Act”.

6       **SEC. 2. FINDINGS.**

7       The Congress finds the following:

8               (1) China has pursued a deliberate, state-di-  
9       rected strategy to dominate global biotechnology, in-

cluding pharmaceutical development, biologics manufacturing, and clinical research and development capabilities.

(2) United States capital flowing to Chinese biotechnology companies through licensing agreements, joint ventures, and equity investments is accelerating China's acquisition of pharmaceutical intellectual property and clinical development capabilities in a manner that creates strategic dependency risks for the United States.

(3) Cross-border out-licensing transactions between United States and European pharmaceutical companies and Chinese biotechnology firms totaled approximately \$136,000,000,000 in 2025, representing a rapid and accelerating transfer of pharmaceutical innovation capacity to entities subject to the direction and control of the People's Republic of China.

(4) Biotechnology, including pharmaceutical development and biologics manufacturing, has civil-military dual-use applications and presents strategic dependency risks for the United States comparable to those presented by semiconductors, artificial intelligence, and other technologies already covered under title VIII of the Defense Production Act of 1950.

1           (5) The BIOSECURE Act, enacted as part of  
2           the National Defense Authorization Act for Fiscal  
3           Year 2026, recognized that biotechnology is both a  
4           national security asset and a strategic vulnerability,  
5           and that the People’s Republic of China seeks to  
6           dominate biotechnology as an industry of the future.

7           (6) Consistent application of outbound invest-  
8           ment screening to biotechnology is necessary to pre-  
9           vent United States capital and intellectual property  
10          from accelerating China’s dominance of the pharma-  
11          ceutical innovation supply chain in a manner that  
12          will create long-term strategic dependency risks  
13          analogous to those the United States now faces in  
14          rare earth elements and semiconductors.

15 **SEC. 3. AMENDMENTS.**

16          Section 809 of the Defense Production Act of 1950  
17          (50 U.S.C. 4589) is amended—

18               (1) in paragraph (10)(A), by adding at the end  
19          the following:

20                       “(vi) Biotechnology, meaning the re-  
21                       search, development, manufacturing, or  
22                       commercialization of—

23                               “(I) pharmaceutical products  
24                               (which has the meaning given the  
25                               term ‘drug’ in section 201(g)(1) of the

1 Federal Food, Drug, and Cosmetic  
2 Act (21 U.S.C. 321(g)(1)));

3 “(II) biological products (as such  
4 term is defined in section 351(i) of  
5 the Public Health Service Act (42  
6 U.S.C. 262(i))); and

7 “(III) therapeutic compounds, in-  
8 cluding drug discovery platforms, clin-  
9 ical research and development capa-  
10 bilities, biologics manufacturing, and  
11 intellectual property and know-how re-  
12 lating to therapeutic compounds,”;

13 (2) in paragraph (7)(A), by adding at the end  
14 the following:

15 “(vi) Biotechnology, meaning the re-  
16 search, development, manufacturing, or  
17 commercialization of—

18 “(I) pharmaceutical products  
19 (which has the meaning given the  
20 term ‘drug’ in section 201(g)(1) of the  
21 Federal Food, Drug, and Cosmetic  
22 Act (21 U.S.C. 321(g)(1)));

23 “(II) biological products (as such  
24 term is defined in section 351(i) of

1 the Public Health Service Act (42  
2 U.S.C. 262(i)); and

3 “(III) therapeutic compounds, in-  
4 cluding drug discovery platforms, clin-  
5 ical research and development capa-  
6 bilities, biologics manufacturing, and  
7 intellectual property and know-how re-  
8 lating to therapeutic compounds,”;  
9 and

10 (3) in paragraph (4)(A), by adding at the end  
11 the following:

12 “(ix) licensing a prohibited technology  
13 from a covered foreign person.”.

14 **SEC. 4. RULEMAKING.**

15 (a) IN GENERAL.—The Secretary of the Treasury  
16 shall, not later than 1 year after the date of the enactment  
17 of this Act, issue a rule to further define the parameters  
18 of the area of “biotechnology” as it is used in paragraphs  
19 (7)(A) and (10)(A) of the Defense Production Act of  
20 1950, as amended by this Act.

21 (b) REQUIREMENTS.—When defining the parameters  
22 of the area of “biotechnology” pursuant to subsection (a),  
23 the Secretary of the Treasury shall—

1           (1) consult with the Secretary of Health and  
2           Human Services, the Secretary of Defense, and the  
3           Director of National Intelligence;

4           (2) give particular consideration to transactions  
5           involving the licensing of intellectual property, drug  
6           discovery platforms, clinical research and develop-  
7           ment capabilities, and biologics manufacturing  
8           know-how to covered foreign persons (as such term  
9           is defined in section 809 of the Defense Production  
10          Act of 1950);

11          (3) give particular consideration to licensing  
12          transactions, joint ventures, and equity investments  
13          involving drug discovery platforms, clinical develop-  
14          ment capabilities, and biologics manufacturing as  
15          priority categories for both the prohibited and  
16          notifiable technology tiers within the biotechnology  
17          sector;

18          (4) consider the degree to which a transaction  
19          would transfer pharmaceutical innovation capacity,  
20          clinical development capabilities, or manufacturing  
21          know-how to entities subject to the direction or con-  
22          trol of the People's Republic of China;

23          (5) define the biotechnology sector to include  
24          the research, development, manufacturing, and com-  
25          mercialization of pharmaceutical products, biological

1 products, and therapeutic compounds, including  
2 drug discovery platforms, clinical research and devel-  
3 opment capabilities, biologics manufacturing, and re-  
4 lated intellectual property and know-how transfers;  
5 and

6 (6) not define the biotechnology sector in a  
7 manner that includes or could be construed to in-  
8 clude agricultural biotechnology, industrial fermenta-  
9 tion unrelated to pharmaceutical or therapeutic pro-  
10 duction, or basic academic research with no direct  
11 pharmaceutical or therapeutic application.

12 **SEC. 5. REPORT REQUIRED.**

13 (a) IN GENERAL.—Not later than 60 days after the  
14 date of the enactment of this Act, the Secretary of Defense  
15 shall submit a report to the appropriate congressional  
16 committees assessing whether flows of United States cap-  
17 ital into China’s biotechnology sector, including through  
18 licensing transactions with Chinese biotechnology firms,  
19 negatively affect United States national security and mili-  
20 tary readiness.

21 (b) FORM.—The report described in subsection (a)  
22 shall be submitted in unclassified form but may include  
23 a classified annex.

1       (c) APPROPRIATE CONGRESSIONAL COMMITTEES DE-  
2 FINED.—The term “appropriate congressional commit-  
3 tees” means—

4           (1) the Committee on Armed Services of the  
5 House of Representatives;

6           (2) the Committee on Financial Services of the  
7 House of Representatives;

8           (3) the Permanent Select Committee on Intel-  
9 ligence of the House of Representatives;

10          (4) the Select Committee on the Strategic Com-  
11 petition between the United States and the Chinese  
12 Communist Party of the House of Representatives;

13          (5) the Committee on Armed Services of the  
14 Senate;

15          (6) the Committee on Banking, Housing, and  
16 Urban Affairs of the Senate; and

17          (7) the Select Committee on Intelligence of the  
18 Senate.

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