

116<sup>TH</sup> CONGRESS  
2<sup>D</sup> SESSION

# H. R. 4866

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## AN ACT

To amend the 21st Century Cures Act to provide for designation of institutions of higher education that provide research, data, and leadership on continuous manufacturing as National Centers of Excellence in Continuous Pharmaceutical Manufacturing, 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 as the “National Centers of  
5 Excellence in Continuous Pharmaceutical Manufacturing  
6 Act of 2020”.

7 **SEC. 2. NATIONAL CENTERS OF EXCELLENCE IN CONTIN-**  
8 **UOUS PHARMACEUTICAL MANUFACTURING.**

9       (a) IN GENERAL.—Section 3016 of the 21st Century  
10 Cures Act (21 U.S.C. 399h) is amended to read as follows:

11 **“SEC. 3016. NATIONAL CENTERS OF EXCELLENCE IN CON-**  
12 **TINUOUS PHARMACEUTICAL MANUFAC-**  
13 **TURING.**

14       “(a) IN GENERAL.—The Secretary of Health and  
15 Human Services, acting through the Commissioner of  
16 Food and Drugs—

17               “(1) shall solicit and, beginning not later than  
18 one year after the date of enactment of the National  
19 Centers of Excellence in Continuous Pharmaceutical  
20 Manufacturing Act of 2020, receive requests from  
21 institutions of higher education to be designated as  
22 a National Center of Excellence in Continuous Phar-  
23 maceutical Manufacturing (in this section referred to  
24 as a ‘National Center of Excellence’) to support the

1 advancement and development of continuous manu-  
2 facturing; and

3 “(2) shall so designate any institution of higher  
4 education that—

5 “(A) requests such designation; and

6 “(B) meets the criteria specified in sub-  
7 section (c).

8 “(b) REQUEST FOR DESIGNATION.—A request for  
9 designation under subsection (a) shall be made to the Sec-  
10 retary at such time, in such manner, and containing such  
11 information as the Secretary may require. Any such re-  
12 quest shall include a description of how the institution of  
13 higher education meets or plans to meet each of the cri-  
14 teria specified in subsection (c).

15 “(c) CRITERIA FOR DESIGNATION DESCRIBED.—The  
16 criteria specified in this subsection with respect to an in-  
17 stitution of higher education are that the institution has,  
18 as of the date of the submission of a request under sub-  
19 section (a) by such institution—

20 “(1) physical and technical capacity for re-  
21 search and development of continuous manufac-  
22 turing;

23 “(2) manufacturing knowledge-sharing net-  
24 works with other institutions of higher education,  
25 large and small pharmaceutical manufacturers, ge-

1       neric and nonprescription manufacturers, contract  
2       manufacturers, and other entities;

3               “(3) proven capacity to design and demonstrate  
4       new, highly effective technology for use in contin-  
5       uous manufacturing;

6               “(4) a track record for creating and transfer-  
7       ring knowledge with respect to continuous manufac-  
8       turing;

9               “(5) the potential to train a future workforce  
10       for research on and implementation of advanced  
11       manufacturing and continuous manufacturing; and

12               “(6) experience in participating in and leading  
13       a continuous manufacturing technology partnership  
14       with other institutions of higher education, large and  
15       small pharmaceutical manufacturers, generic and  
16       nonprescription manufacturers, contract manufac-  
17       turers, and other entities—

18                       “(A) to support companies with continuous  
19       manufacturing in the United States;

20                       “(B) to support Federal agencies with  
21       technical assistance, which may include regu-  
22       latory and quality metric guidance as applica-  
23       ble, for advanced manufacturing and continuous  
24       manufacturing;

1           “(C) with respect to continuous manufac-  
2           turing, to organize and conduct research and  
3           development activities needed to create new and  
4           more effective technology, capture and dissemi-  
5           nate expertise, create intellectual property, and  
6           maintain technological leadership;

7           “(D) to develop best practices for design-  
8           ing continuous manufacturing; and

9           “(E) to assess and respond to the work-  
10          force needs for continuous manufacturing, in-  
11          cluding the development of training programs if  
12          needed.

13          “(d) TERMINATION OF DESIGNATION.—The Sec-  
14          retary may terminate the designation of any National Cen-  
15          ter of Excellence designated under this section if the Sec-  
16          retary determines such National Center of Excellence no  
17          longer meets the criteria specified in subsection (c). Not  
18          later than 60 days before the effective date of such a ter-  
19          mination, the Secretary shall provide written notice to the  
20          National Center of Excellence, including the rationale for  
21          such termination.

22          “(e) CONDITIONS FOR DESIGNATION.—As a condi-  
23          tion of designation as a National Center of Excellence  
24          under this section, the Secretary shall require that an in-

1 stitution of higher education enter into an agreement with  
2 the Secretary under which the institution agrees—

3 “(1) to collaborate directly with the Food and  
4 Drug Administration to publish the reports required  
5 by subsection (g);

6 “(2) to share data with the Food and Drug Ad-  
7 ministration regarding best practices and research  
8 generated through the funding under subsection (f);

9 “(3) to develop, along with industry partners  
10 (which may include large and small biopharma-  
11 ceutical manufacturers, generic and nonprescription  
12 manufacturers, and contract manufacturers) and an-  
13 other institution or institutions designated under  
14 this section, if any, a roadmap for developing a con-  
15 tinuous manufacturing workforce;

16 “(4) to develop, along with industry partners  
17 and other institutions designated under this section,  
18 a roadmap for strengthening existing, and devel-  
19 oping new, relationships with other institutions; and

20 “(5) to provide an annual report to the Food  
21 and Drug Administration regarding the institution’s  
22 activities under this section, including a description  
23 of how the institution continues to meet and make  
24 progress on the criteria listed in subsection (c).

25 “(f) FUNDING.—

1           “(1) IN GENERAL.—The Secretary shall award  
2 funding, through grants, contracts, or cooperative  
3 agreements, to the National Centers of Excellence  
4 designated under this section for the purpose of  
5 studying and recommending improvements to contin-  
6 uous manufacturing, including such improvements  
7 as may enable the Centers—

8           “(A) to continue to meet the conditions  
9 specified in subsection (e); and

10           “(B) to expand capacity for research on,  
11 and development of, continuing manufacturing.

12           “(2) CONSISTENCY WITH FDA MISSION.—As a  
13 condition on receipt of funding under this sub-  
14 section, a National Center of Excellence shall agree  
15 to consider any input from the Secretary regarding  
16 the use of funding that would—

17           “(A) help to further the advancement of  
18 continuous manufacturing through the National  
19 Center of Excellence; and

20           “(B) be relevant to the mission of the  
21 Food and Drug Administration.

22           “(3) AUTHORIZATION OF APPROPRIATIONS.—  
23 There is authorized to be appropriated to carry out  
24 this subsection \$80,000,000 for the period of fiscal  
25 years 2021 through 2025.

1           “(4) RULE OF CONSTRUCTION.—Nothing in  
2 this section shall be construed as precluding a Na-  
3 tional Center for Excellence designated under this  
4 section from receiving funds under any other provi-  
5 sion of this Act or any other Federal law.

6           “(g) ANNUAL REVIEW AND REPORTS.—

7           “(1) ANNUAL REPORT.—Beginning not later  
8 than one year after the date on which the first des-  
9 ignation is made under subsection (a), and annually  
10 thereafter, the Secretary shall—

11                   “(A) submit to Congress a report describ-  
12 ing the activities, partnerships and collabora-  
13 tions, Federal policy recommendations, previous  
14 and continuing funding, and findings of, and  
15 any other applicable information from, the Na-  
16 tional Centers of Excellence designated under  
17 this section; and

18                   “(B) make such report available to the  
19 public in an easily accessible electronic format  
20 on the website of the Food and Drug Adminis-  
21 tration.

22           “(2) REVIEW OF NATIONAL CENTERS OF EX-  
23 CELLENCE AND POTENTIAL DESIGNEES.—The Sec-  
24 retary shall periodically review the National Centers  
25 of Excellence designated under this section to ensure



1 that such National Centers of Excellence continue to  
2 meet the criteria for designation under this section.

3 “(3) REPORT ON LONG-TERM VISION OF FDA  
4 ROLE.—Not later than 2 years after the date on  
5 which the first designation is made under subsection  
6 (a), the Secretary, in consultation with the National  
7 Centers of Excellence designated under this section,  
8 shall submit a report to the Congress on the long-  
9 term vision of the Department of Health and  
10 Human Services on the role of the Food and Drug  
11 Administration in supporting continuous manufac-  
12 turing, including—

13 “(A) a national framework of principles re-  
14 lated to the implementation and regulation of  
15 continuous manufacturing;

16 “(B) a plan for the development of Federal  
17 regulations and guidance for how advanced  
18 manufacturing and continuous manufacturing  
19 can be incorporated into the development of  
20 pharmaceuticals and regulatory responsibilities  
21 of the Food and Drug Administration; and

22 “(C) appropriate feedback solicited from  
23 the public, which may include other institutions,  
24 large and small biopharmaceutical manufactur-

1           ers, generic and nonprescription manufacturers,  
2           and contract manufacturers.

3           “(h) DEFINITIONS.—In this section:

4           “(1) ADVANCED MANUFACTURING.—The term  
5           ‘advanced manufacturing’ means an approach for  
6           the manufacturing of pharmaceuticals that incor-  
7           porates novel technology, or uses an established  
8           technique or technology in a new or innovative way  
9           (such as continuous manufacturing where the input  
10          materials are continuously transformed within the  
11          process by two or more unit operations) that en-  
12          hances drug quality or improves the manufacturing  
13          process.

14          “(2) CONTINUOUS MANUFACTURING.—The  
15          term ‘continuous manufacturing’—

16                  “(A) means a process where the input ma-  
17                  terials are continuously fed into and trans-  
18                  formed within the process, and the processed  
19                  output materials are continuously removed from  
20                  the system; and

21                  “(B) consists of an integrated process that  
22                  consists of a series of two or more unit oper-  
23                  ations.

24          “(3) INSTITUTION OF HIGHER EDUCATION.—  
25          The term ‘institution of higher education’ has the

1 meaning given such term in section 101(a) of the  
2 Higher Education Act of 1965 (20 U.S.C. 1001(a)).

3 “(4) SECRETARY.—The term ‘Secretary’ means  
4 the Secretary of Health and Human Services, acting  
5 through the Commissioner of Food and Drugs.”.

6 (b) TRANSITION RULE.—Section 3016 of the 21st  
7 Century Cures Act (21 U.S.C. 399h), as in effect on the  
8 day before the date of the enactment of this section, shall  
9 apply with respect to grants awarded under such section  
10 before such date of enactment.

Passed the House of Representatives September 21,  
2020.

Attest:

*Clerk.*

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