

117TH CONGRESS  
1ST SESSION

# H. R. 3932

To establish a program to develop antimicrobial innovations targeting the most challenging pathogens and most threatening infections.

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## IN THE HOUSE OF REPRESENTATIVES

JUNE 16, 2021

Mr. MICHAEL F. DOYLE of Pennsylvania (for himself and Mr. FERGUSON) introduced the following bill; which was referred to the Committee on Energy and Commerce, and in addition to the Committees on Ways and Means, Veterans' Affairs, Armed Services, the Judiciary, and Homeland Security, for a period to be subsequently determined by the Speaker, in each case for consideration of such provisions as fall within the jurisdiction of the committee concerned

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## A BILL

To establish a program to develop antimicrobial innovations targeting the most challenging pathogens and most threatening infections.

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 “Pioneering Anti-  
5 microbial Subscriptions To End Up surging Resistance  
6 Act of 2021” or the “PASTEUR Act of 2021”.

1 **SEC. 2. DEVELOPING ANTIMICROBIAL INNOVATIONS.**

2 Title III of the Public Health Service Act (42 U.S.C.  
3 241 et seq.) is amended by adding at the end the fol-  
4 lowing:

5 **“PART W—DEVELOPING ANTIMICROBIAL**  
6 **INNOVATIONS**

7 **“SEC. 3990O. ESTABLISHMENT OF COMMITTEE; SUBSCRIP-**  
8 **TION MODEL; ADVISORY GROUP.**

9 “(a) IN GENERAL.—Not later than 60 days after the  
10 date of enactment of this part, the Secretary shall estab-  
11 lish a Committee on Critical Need Antimicrobials and ap-  
12 point members to the Committee.

13 “(b) MEMBERS.—

14 “(1) IN GENERAL.—The Committee shall con-  
15 sist of at least one representative from each of the  
16 National Institute of Allergy and Infectious Dis-  
17 eases, the Centers for Disease Control and Preven-  
18 tion, the Biomedical Advanced Research and Devel-  
19 opment Authority, the Food and Drug Administra-  
20 tion, the Centers for Medicare & Medicaid Services,  
21 the Veterans Health Administration, and the De-  
22 partment of Defense.

23 “(2) CHAIR.—The Secretary shall appoint one  
24 of the members of the Committee to serve as the  
25 Chair of the Committee.

1       “(c) DUTIES.—Not later than 1 year after the ap-  
2 pointment of all initial members of the Committee, the  
3 Secretary, in collaboration with the Committee, and in  
4 consultation with the Critical Need Antimicrobials Advi-  
5 sory Group established under subsection (g), shall do the  
6 following:

7               “(1) Develop a list of infections for which new  
8 antimicrobial drug development is needed, taking  
9 into account organisms, sites of infection, and type  
10 of infections for which there is an unmet medical  
11 need, findings from the most recent report entitled  
12 ‘Antibiotic Resistance Threats in the United States’  
13 issued by the Centers for Disease Control and Pre-  
14 vention, or an anticipated unmet medical need, in-  
15 cluding a potential global health security threat. For  
16 the list developed under this paragraph, the Sec-  
17 retary, in collaboration with the Committee, may use  
18 the infection list in such most recent report for up  
19 to 3 years following the date of enactment of this  
20 part and subsequently update the list under this  
21 paragraph in accordance with subsection (e).

22               “(2) Develop regulations, in accordance with  
23 subsection (d), outlining favored characteristics of  
24 critical need antimicrobial drugs, that are evidence  
25 based, clinically focused, and designed to treat the

1 infections described in paragraph (1), and estab-  
2 lishing criteria for how each such characteristic will  
3 adjust the monetary value of a subscription contract  
4 awarded under subsection (f) or section 399QQ. The  
5 favored characteristics shall be weighed for purposes  
6 of such monetary value such that meeting certain  
7 characteristics, or meeting more than one such char-  
8 acteristic, increases the monetary value. Such fa-  
9 vored characteristics of an antimicrobial drug shall  
10 include—

11 “(A) treating infections on the list under  
12 paragraph (1);

13 “(B) improving clinical outcomes for pa-  
14 tients with multi-drug-resistant infections;

15 “(C) being a first-approved antimicrobial  
16 drug that has the potential to address unmet  
17 medical needs for the treatment of a serious or  
18 life-threatening infection, and, to a lesser ex-  
19 tent, second and third drugs that treat such in-  
20 fections;

21 “(D) route of administration, especially  
22 through oral administration;

23 “(E)(i) containing no active moiety (as de-  
24 fined by the Secretary in section 314.3 of title  
25 21, Code of Federal Regulations (or any suc-

1           cessor regulations)) that has been approved in  
2           any other application under section 505(b) of  
3           the Federal Food, Drug, and Cosmetic Act or  
4           intending to be the subject of a new original  
5           biologics license application under section  
6           351(a);

7           “(ii) being a member of a new class of  
8           drugs with a novel target and novel mode of ac-  
9           tion that are distinctly different from the target  
10          or mode of any antimicrobial drug approved  
11          under section 505 of such Act or licensed under  
12          section 351, including reduced toxicity;

13          “(iii) not being affected by cross-resistance  
14          to any antimicrobial drug approved under such  
15          section 505 or licensed under such section 351;

16          “(F) addressing a multi-drug resistant in-  
17          fection through a novel chemical scaffold or  
18          mechanism of action;

19          “(G) having received a transitional sub-  
20          scription contract under subsection (f); and

21          “(H) any other characteristic the Sec-  
22          retary, in collaboration with the Committee, de-  
23          termines necessary.

24          “(d) REGULATIONS.—

1           “(1) IN GENERAL.—Not later than 1 year after  
2 the appointment of the initial members of the Com-  
3 mittee, the Secretary shall issue proposed regula-  
4 tions which shall include—

5           “(A) a process by which the sponsors can  
6 apply for an antimicrobial drug to become a  
7 critical need antimicrobial drug under section  
8 399PP;

9           “(B) how subscription contracts under  
10 such section shall be established and paid;

11           “(C) the favored characteristics under sub-  
12 section (c)(2), how such characteristics will be  
13 weighed, and the minimum number and kind of  
14 favored characteristics needed for an anti-  
15 microbial drug to be designated a critical need  
16 antimicrobial drug; and

17           “(D) other elements of the subscription  
18 contract process, in accordance with this part.

19           “(2) DEVELOPMENT OF FINAL REGULA-  
20 TIONS.—Before finalizing the regulations under  
21 paragraph (1), the Secretary shall solicit public com-  
22 ment and hold public meetings for the period begin-  
23 ning on the date on which the proposed regulations  
24 are issued and ending on the date that is 120 days  
25 after such date of issuance. The Secretary shall fi-

1       nalize and publish such regulations not later than  
2       120 days after the close of such period of public  
3       comment and meetings.

4               “(3) SUBSCRIPTION CONTRACT OFFICE.—Not  
5       later than 6 months after the date of enactment of  
6       this part, the Secretary shall propose an agency or  
7       office in the Department of Health and Human  
8       Services to manage the establishment and payment  
9       of subscription contracts awarded under section  
10      399QQ, including eligibility, requirements, and con-  
11      tract amounts. The Secretary shall solicit public  
12      comment and finalize the agency or office no later  
13      than 45 days following the proposed agency or of-  
14      fice. Such agency or office shall be referred to as the  
15      ‘Subscription Contract Office’.

16              “(e) LIST OF INFECTIONS.—The Secretary, in col-  
17      laboration with the Committee, shall update the list of in-  
18      fections under subsection (c)(1) at least every 2 years.

19              “(f) TRANSITIONAL SUBSCRIPTION CONTRACTS.—

20                      “(1) IN GENERAL.—Not earlier than 30 days  
21      after the date of enactment of this part and ending  
22      on the date that the Secretary finalizes the subscrip-  
23      tion contract regulations under subsection (d), the  
24      Secretary may use up to \$1,000,000,000 of the  
25      amount appropriated under section 399SS(a) to en-

1 gage in transitional subscription contracts of up to  
2 3 years in length with antimicrobial developers, as  
3 determined by the Secretary, that have developed  
4 antimicrobial drugs treating infections listed in the  
5 most recent report entitled ‘Antibiotic Resistance  
6 Threats in the United States’ issued by the Centers  
7 for Disease Control and Prevention, and may include  
8 antimicrobial drugs that are qualified infectious dis-  
9 ease products (as defined in section 505E(g) of the  
10 Federal Food, Drug, and Cosmetic Act), innovative  
11 biological products, or innovative drugs that achieve  
12 a clinical outcome through immunomodulation. Such  
13 a contract may authorize the contractor to use funds  
14 made available under the contract for completion of  
15 postmarketing clinical studies, manufacturing, and  
16 other preclinical and clinical efforts.

17 “(2) REQUIREMENTS.—

18 “(A) IN GENERAL.—The Secretary,  
19 through the office described in paragraph (4),  
20 may enter into a contract under paragraph  
21 (1)—

22 “(i) if the Secretary determines that  
23 the antimicrobial drug is intended to treat  
24 an infection for which there is an unmet



1 clinical need, an anticipated clinical need,  
2 or drug resistance;

3 “(ii) subject to terms including—

4 “(I) that the Secretary shall  
5 cease any payment installments under  
6 a transitional subscription contract if  
7 the sponsor does not—

8 “(aa) ensure commercial and  
9 Federal availability of the anti-  
10 microbial drug within 30 days of  
11 receiving first payment under the  
12 contract;

13 “(bb) identify, track, and  
14 publicly report drug resistance  
15 data and trends using available  
16 data related to the antimicrobial  
17 drug;

18 “(cc) develop and implement  
19 education and communications  
20 strategies, including communica-  
21 tions for individuals with limited  
22 English proficiency and individ-  
23 uals with disabilities, for health  
24 care professionals and patients

1 about appropriate use of the  
2 antimicrobial drug;

3 “(dd) submit a plan for reg-  
4 istering the antimicrobial drug in  
5 additional countries where an  
6 unmet medical need exists, which  
7 such plan may be consistent with  
8 the Stewardship and Access Plan  
9 (SAP) Development Guide  
10 (2021);

11 “(ee) subject to subpara-  
12 graph (B), ensure a reliable drug  
13 supply chain, thus leading to an  
14 interruption of the supply of the  
15 antimicrobial drug in the United  
16 States for more than 60 days; or

17 “(ff) make meaningful  
18 progress toward completion of  
19 Food and Drug Administration-  
20 required postmarketing studies,  
21 including such studies that are  
22 evidence based; and

23 “(II) other terms as determined  
24 by the Secretary; and

25 “(iii) if—

1                   “(I) a phase 3 clinical study has  
2                   been initiated for the antimicrobial  
3                   drug; or

4                   “(II) the antimicrobial drug has  
5                   been approved under section 505(c) of  
6                   the Federal Food, Drug, and Cos-  
7                   metic Act or licensed under section  
8                   351(a).

9                   “(B) WAIVER.—The requirement under  
10                  subparagraph (A)(ii)(I)(ee) may be waived in  
11                  the case that an emergency prohibits access to  
12                  a reliable drug supply chain.

13                  “(3) TRANSITIONAL GUIDANCE.—Not later  
14                  than 120 days after the appointment of the initial  
15                  members of the Committee, the Secretary shall  
16                  issue, in consultation with the Committee, transi-  
17                  tional guidance outlining the antimicrobial drugs  
18                  that are eligible for transitional subscription con-  
19                  tracts under paragraph (1), the requirements to  
20                  enter into a transitional subscription contract under  
21                  paragraph (2), and the process by which drug devel-  
22                  opers can enter into transitional subscription con-  
23                  tracts with the Secretary under this subsection.

24                  “(4) PAYMENT OFFICE AND MECHANISM.—Not  
25                  later than 30 days after the date of enactment of

1 this part, the Secretary shall determine the agency  
2 or office in the Department of Health and Human  
3 Services that will manage the transitional subscrip-  
4 tion contracts, including eligibility, requirements,  
5 and contract amounts, during the period described  
6 in paragraph (1).

7 “(g) CRITICAL NEED ANTIMICROBIAL ADVISORY  
8 GROUP.—

9 “(1) IN GENERAL.—Not later than 30 days  
10 after the appointment of all initial members of the  
11 Committee, the Secretary, in collaboration with the  
12 Committee, shall establish a Critical Need Anti-  
13 microbial Advisory Group (referred to in this sub-  
14 section as the ‘Advisory Group’) and appoint mem-  
15 bers to the Advisory Group.

16 “(2) MEMBERS.—The members of the Advisory  
17 Group shall include—

18 “(A) not fewer than 6 individuals who  
19 are—

20 “(i) infectious disease specialists; or

21 “(ii) other health experts with exper-  
22 tise in researching antimicrobial resistance,  
23 health economics, or commercializing anti-  
24 microbial drugs; and

25 “(B) not fewer than 5 patient advocates.

1           “(3) CHAIR.—The Secretary shall appoint one  
2 of the members of the Advisory Group to serve as  
3 the Chair.

4           “(4) CONFLICTS OF INTEREST.—In appointing  
5 members under paragraph (2), the Secretary shall  
6 ensure that no member receives compensation in any  
7 manner from a commercial or for-profit entity that  
8 develops antimicrobials or that might benefit from  
9 antimicrobial development.

10           “(5) APPLICABILITY OF FACa.—Except as oth-  
11 erwise provided in this subsection, the Federal Advi-  
12 sory Committee Act shall apply to the Advisory  
13 Group.

14 **“SEC. 399PP. CRITICAL NEED ANTIMICROBIAL DRUG APPLI-**  
15 **CATION AND PAYMENT THROUGH SUBSCRIP-**  
16 **TION CONTRACTS.**

17           “(a) IN GENERAL.—

18           “(1) SUBMISSION OF REQUEST.—The sponsor  
19 of an application under section 505(b) of the Fed-  
20 eral Food, Drug, and Cosmetic Act or section 351(a)  
21 for an antimicrobial drug may request that the Sec-  
22 retary designate the drug as a critical need anti-  
23 microbial. A request for such designation may be  
24 submitted after the Secretary grants for such drug  
25 an investigational new drug exemption under section

1 505(i) of the Federal Food, Drug, and Cosmetic Act  
2 or section 351(a)(3), and shall be submitted not  
3 later than 5 years after the date of approval under  
4 section 505(c) of the Federal Food, Drug, and Cos-  
5 metic Act or licensure under section 351(a).

6 “(2) CONTENT OF REQUEST.—A request under  
7 paragraph (1) shall include information, such as  
8 clinical, preclinical and postmarketing data, a list of  
9 the favorable characteristics described in section  
10 39900(c)(2), and any other material that the Sec-  
11 retary in consultation with the Committee requires.

12 “(3) REVIEW BY SECRETARY.—The Secretary  
13 shall promptly review all requests for designation  
14 submitted under this subsection, assess all required  
15 application components, and determine if the anti-  
16 microbial drug is likely to meet the favorable charac-  
17 teristics identified in the application upon the com-  
18 pletion of clinical development. After review, the Sec-  
19 retary shall approve or deny each request for des-  
20 ignation not later than 90 days after receiving a re-  
21 quest. If the Secretary approves a request, it shall  
22 publish the value of the contract that the critical  
23 need antimicrobial developer would be eligible to re-  
24 ceive if such developer successfully demonstrates

1 that the drug meets the maximum value of the fa-  
2 vored characteristics listed in the application.

3 “(4) LENGTH OF DESIGNATION PERIOD.—A  
4 designation granted under this section shall be in ef-  
5 fect for a period of 10 years after the date that the  
6 designation is approved, and shall remain in effect  
7 for such period even if the infection treated by such  
8 drug is later removed from the list of infections  
9 under section 39900(c)(1).

10 “(5) SUBSEQUENT REVIEWS.—No sooner than  
11 2 years after a designation approval or denial under  
12 subsection (3), the sponsor may request a subse-  
13 quent review to re-evaluate the value of a contract  
14 to include any new information.

15 “(b) DEVELOPMENT OF DESIGNATED DRUGS.—If a  
16 critical need antimicrobial designation is granted during  
17 clinical development of an antimicrobial drug, the Sec-  
18 retary may work with the sponsor to maximize the oppor-  
19 tunity for the sponsor to successfully demonstrate that the  
20 antimicrobial drug possesses the favored characteristics of  
21 high-monetary valued products identified under section  
22 39900(c)(2).

23 “(c) APPROPRIATE USE OF CRITICAL NEED ANTI-  
24 MICROBIAL.—

1           “(1) IN GENERAL.—The sponsor of an anti-  
2           microbial drug that receives designation under sub-  
3           section (a) shall within 90 days of such designation,  
4           submit to the Secretary a plan for appropriate use  
5           of diagnostics, in order for the Secretary and Com-  
6           mittee to consider such plan in developing clinical  
7           guidelines. An appropriate use plan—

8                   “(A) shall include—

9                           “(i) the appropriate use of the drug;

10                           and

11                           “(ii) the appropriate use of diagnostic  
12                           tools, where available, such as diagnostic  
13                           testing for biomarkers related to anti-  
14                           microbial-resistant pathogens, or other tar-  
15                           geted diagnostic approaches, to inform use  
16                           of the drug; and

17                   “(B) may be developed in partnership with  
18           the Secretary, infectious disease experts, diag-  
19           nostic experts or developers, laboratory experts,  
20           or another entity.

21           “(2) CONSULTATION.—The Secretary shall con-  
22           sult with relevant professional societies and the Crit-  
23           ical Need Antimicrobial Advisory Group established  
24           under section 39900(g) to ensure that clinical  
25           guidelines issued by the Secretary under paragraph



1 (3), with respect to an antimicrobial drug designated  
2 under subsection (a), includes the use of appropriate  
3 diagnostic approaches, taking into consideration the  
4 diagnostic plan submitted by a sponsor under para-  
5 graph (1).

6 “(3) PUBLICATION OF CLINICAL GUIDELINES.—  
7 Not later than 1 year after the Secretary makes the  
8 first designation under subsection (a), and not less  
9 than every 3 years thereafter, the Secretary shall  
10 publish clinical guidelines in consultation with rel-  
11 evant professional societies with respect to each anti-  
12 microbial drug that has been approved or licensed as  
13 described in subsection (a)(1) and that has been des-  
14 ignated under subsection (a), which guidelines shall  
15 set forth the evidence-based recommendations for  
16 prescribing the drug, in accordance with the submis-  
17 sions of the sponsor under paragraph (1) and after  
18 consultation under paragraph (2), as appropriate.

19 **“SEC. 399QQ. SUBSCRIPTION CONTRACTS.**

20 “(a) APPLICATION FOR A SUBSCRIPTION CON-  
21 TRACT.—

22 “(1) SUBMISSION OF APPLICATIONS.—After ap-  
23 proval under section 505(c) of the Federal Food,  
24 Drug, and Cosmetic Act or licensure under section  
25 351(a), the sponsor of an antimicrobial drug des-

1       ignated as a critical need antimicrobial under section  
2       399PP may submit an application for a subscription  
3       contract with the Secretary, under a procedure es-  
4       tablished by the Secretary.

5               “(2) REVIEW OF APPLICATIONS.—The Sec-  
6       retary shall, in consultation with the Committee—

7                       “(A) review all applications for subscrip-  
8       tion contracts under paragraph (1) and assess  
9       all required application components;

10                      “(B) determine the extent to which the  
11       critical need antimicrobial meets the favored  
12       characteristics identified under section  
13       399OO(c)(2), and deny any application for a  
14       drug that meets none of such characteristics;  
15       and

16                      “(C) assign a monetary value to the con-  
17       tract based on the regulations developed under  
18       section 399OO(d).

19       “(b) CRITERIA.—To qualify for a subscription con-  
20       tract under this section, the sponsor of an antimicrobial  
21       drug designated as a critical need antimicrobial shall agree  
22       to—

23                      “(1) ensure commercial and Federal availability  
24       of the antimicrobial drug within 30 days of receiving

1 first payment under the contract, and sufficient sup-  
2 ply for susceptibility device manufacturers;

3 “(2) identify, track, and publicly report drug  
4 resistance data and trends using available data re-  
5 lated to the antimicrobial drug;

6 “(3) develop and implement education and com-  
7 munications strategies, including communications  
8 for individuals with limited English proficiency and  
9 individuals with disabilities, for health care profes-  
10 sionals and patients about appropriate use of the  
11 antimicrobial drug;

12 “(4) submit an appropriate use assessment to  
13 the Secretary, Committee, Food and Drug Adminis-  
14 tration, and Centers for Disease Control and Pre-  
15 vention every 2 years regarding use of the anti-  
16 microbial drug, including how the drug is being mar-  
17 keted;

18 “(5) submit a plan for registering the drug in  
19 additional countries where an unmet medical need  
20 exists;

21 “(6) ensure a reliable drug supply chain, where  
22 any interruption to the supply chain will not last for  
23 more than 60 days in the United States;

1           “(7) complete any postmarketing studies re-  
2           quired by the Food and Drug Administration in a  
3           timely manner;

4           “(8) produce the drug at a reasonable volume  
5           determined with the Secretary to ensure patient ac-  
6           cess to the drug;

7           “(9) price the drug at a price that is not lower  
8           than a comparable generic drug;

9           “(10) abide by the manufacturing and environ-  
10          mental best practices in the supply chain to ensure  
11          that there is no discharge into, or contamination of,  
12          the environment by antimicrobial agents or products  
13          as a result of the manufacturing process; and

14          “(11) abide by other terms as the Secretary  
15          may require.

16          “(c) AMOUNT AND TERMS OF CONTRACTS.—

17                 “(1) AMOUNTS.—A subscription contract under  
18                 this section shall be for the sale to the Secretary of  
19                 any quantity of the antimicrobial drug needed over  
20                 the term of the contract under paragraph (2), at an  
21                 agreed upon price, for a total projected amount de-  
22                 termined by the Secretary that is not less than  
23                 \$750,000,000 and not more than \$3,000,000,000,  
24                 adjusted for inflation, accounting for the favored  
25                 characteristics of the drug, as determined by the

1 Secretary, in consultation with the Committee, under  
2 subsection (a)(2), and shall be allocated from the  
3 amount made available under section 399SS(a). Not  
4 later than 6 months after the subscription contract  
5 is granted under subsection (a), the Secretary shall  
6 provide payments for purchased drugs in install-  
7 ments established by the Secretary in consultation  
8 with the sponsor of the antimicrobial drug and in ac-  
9 cordance with subsection (d)(3). Funds received by  
10 the sponsor shall be used to support criteria quali-  
11 fication under subsection (b), the completion of post-  
12 marketing clinical studies, manufacturing, other pre-  
13 clinical and clinical activities, or other activities  
14 agreed to by the Secretary and sponsor in the con-  
15 tract.

16 “(2) TERMS.—

17 “(A) INITIAL TERM.—The initial term of a  
18 contract under this subsection shall be no less  
19 than 5 years or greater than the greater of 10  
20 years or the remaining period of time during  
21 which the sponsor has patent protections or a  
22 remaining exclusivity period with respect to the  
23 antimicrobial drug in the United States, as list-  
24 ed in the publication of the Food and Drug Ad-  
25 ministration entitled ‘Approved Drug Products

1 with Therapeutic Equivalence Evaluations’.  
2 Payments may be in equal annual installments  
3 with the option to redeem 50 percent of the last  
4 year’s reimbursement in year 1 of the contract  
5 in order to offset costs of establishing manufac-  
6 turing capacity, or another subscription ar-  
7 rangement to which the Secretary and sponsor  
8 agree. Subscription contracts shall remain in ef-  
9 fect for such period even if the infection treated  
10 by such antimicrobial drug is later removed  
11 from the list of infections under section  
12 39900(c)(1).

13 “(B) EXTENSION OF CONTRACTS.—The  
14 Secretary may extend a subscription contract  
15 with a sponsor under this subsection beyond the  
16 initial contract period. A single contract exten-  
17 sion may be in effect not later than the date on  
18 which all periods of exclusivity granted by the  
19 Food and Drug Administration expire and shall  
20 be in an amount not to exceed \$25,000,000 per  
21 year. All other terms of an extended contract  
22 shall be the same as the terms of the initial  
23 contract. The total amount of funding used on  
24 such contract extensions shall be no more than

1           \$1,000,000,000, and shall be allocated from the  
2           amount made available under section 399SS.

3           “(C) MODIFICATION OF CONTRACTS.—The  
4           Secretary or sponsor, 1 year after the start of  
5           the contract period under this subsection and  
6           every 2 years thereafter, may request a modi-  
7           fication of the amount of the contract based on  
8           information that adjusts favored characteristics  
9           in section 399OO(c)(2).

10          “(3) ADJUSTMENT.—In the case of an anti-  
11          microbial drug that received a transitional subscrip-  
12          tion contract under section 399OO(f), the amount of  
13          a subscription contract for such drug under this sec-  
14          tion shall be reduced by the amount of the transi-  
15          tional subscription contract under such section  
16          399OO(f) for such drug.

17          “(4) CONTRACTS FOR GENERIC AND BIO-  
18          SIMILAR VERSIONS.—Notwithstanding any other  
19          provision in this part, the Secretary may award a  
20          subscription contract under this section to a manu-  
21          facturer of a generic or biosimilar version of an anti-  
22          microbial drug for which a subscription contract has  
23          been awarded under this section. Such contracts  
24          shall be awarded in accordance with a procedure, in-

1 including for determining the terms and amounts of  
2 such contracts, established by the Secretary.

3 “(d) ANNUAL ANTIMICROBIAL DRUG SPONSOR REV-  
4 ENUE LIMITATIONS.—

5 “(1) REPORTING REQUIREMENT.—

6 “(A) IN GENERAL.—Not later than a date  
7 determined appropriate by the Secretary fol-  
8 lowing the end of each calendar year, and not  
9 earlier than 6 months after the end of each cal-  
10 endar year, the head (or a designee of such  
11 head) of each Federal agency carrying out a  
12 specified government program shall, in accord-  
13 ance with this paragraph, report to the Sub-  
14 scription Contract Office established under sec-  
15 tion 3990O(d)(3) the total prescription drug  
16 sales for each applicable antimicrobial drug  
17 under contract with respect to such program for  
18 such calendar year.

19 “(B) MEDICARE PART D PROGRAM.—For  
20 purposes of subparagraph (A), the Secretary  
21 shall report, for each applicable antimicrobial  
22 drug covered under part D of title XVIII of the  
23 Social Security Act, the product of—

24 “(i) the per-unit ingredient cost, as  
25 reported to the Secretary by prescription



1 drug plans and Medicare Advantage pre-  
2 scription drug plans, minus any per-unit  
3 rebate, discount, or other price concession  
4 provided by the sponsor of such applicable  
5 antimicrobial drug, as reported to the Sec-  
6 retary by the prescription drug plans and  
7 the Medicare Advantage prescription drug  
8 plans; and

9 “(ii) the number of units of such ap-  
10 plicable antimicrobial drug paid for under  
11 such part D.

12 “(C) MEDICARE PART B PROGRAM.—

13 “(i) IN GENERAL.—For purposes of  
14 subparagraph (A), the Secretary shall re-  
15 port, for each applicable antimicrobial drug  
16 covered under part B of title XVIII of the  
17 Social Security Act, the product of—

18 “(I) the per-unit average sales  
19 price (as defined in section 1847A(c)  
20 of such Act) or the per-unit payment  
21 rate under such part B for a sepa-  
22 rately paid prescription drug without  
23 a reported average sales price; and

1                   “(II) the number of units of such  
2                   applicable antimicrobial drug paid for  
3                   under such part B.

4                   “(ii) UNITS AND ALLOCATED  
5                   PRICES.—The Secretary shall establish a  
6                   process for determining the units and the  
7                   allocated price for purposes of this sub-  
8                   paragraph for those applicable anti-  
9                   microbial drugs that are not separately  
10                  payable or for which National Drug Codes  
11                  are not reported.

12                  “(D) MEDICARE PART A PROGRAM.—

13                  “(i) IN GENERAL.—For purposes of  
14                  subparagraph (A), the Secretary shall re-  
15                  port, for each applicable antimicrobial drug  
16                  covered under part A of title XVIII of the  
17                  Social Security Act, the product of—

18                         “(I) the per-unit price under  
19                         such part A for the antimicrobial  
20                         drug; and

21                         “(II) the number of units of such  
22                         antimicrobial drug paid for under  
23                         such part A.

24                         “(ii) SPECIAL RULE.—For purposes of  
25                         clause (i), the Secretary shall establish a

1 process for determining the units and the  
2 allocated price for those prescription drugs  
3 that are not separately payable or for  
4 which National Drug Codes are not re-  
5 ported in the diagnosis-related groups.

6 “(E) MEDICAID PROGRAM.—Under the au-  
7 thority of section 1902(a)(6) of the Social Secu-  
8 rity Act, the Secretary shall require each State  
9 that makes medical assistance available under  
10 the State plan under title XIX of such Act (or  
11 any waiver of such plan) for an applicable anti-  
12 microbial drug (including, if applicable, any  
13 such drug which is a covered outpatient drug  
14 under a rebate agreement entered into under  
15 section 1927 of such Act) to report, in a form  
16 consistent with a standard reporting format es-  
17 tablished by the Secretary, not later than the  
18 date determined under subparagraph (A)—

19 “(i) information on the total number  
20 of units of each dosage form and strength  
21 and package size of each applicable anti-  
22 microbial drug dispensed during the pre-  
23 ceding calendar year under such State plan  
24 or waiver (including any such drugs dis-  
25 pensed to an individual enrolled with a

1           medicaid managed care organization or  
2           other specified entity (as such terms are  
3           defined in section 1903(m) of such Act));  
4           and

5           “(ii) with respect to each dosage form  
6           and strength and package size of each such  
7           drug, the amount equal to—

8                   “(I) the product of—

9                           “(aa) the total number of  
10                           units dispensed under the State  
11                           plan or waiver during the pre-  
12                           ceding calendar year (as deter-  
13                           mined under clause (i)); and

14                           “(bb) the per-unit ingredient  
15                           cost paid by the State for each  
16                           such unit; minus

17                   “(II) any discounts or other price  
18                   concessions provided and rebates paid  
19                   to the State with respect to the dos-  
20                   age form and strength and package  
21                   size of such drug and such calendar  
22                   year (including rebates paid under a  
23                   rebate agreement under section 1927  
24                   of such Act and any State supple-

1                   mental rebates paid under a supple-  
2                   mental rebate agreement).

3                   “(F) DEPARTMENT OF VETERANS AF-  
4                   FAIRS.—For purposes of subparagraph (A), the  
5                   Secretary of Veterans Affairs shall report the  
6                   total amount paid for each applicable anti-  
7                   microbial drug procured by the Veterans Health  
8                   Administration for individuals who receive  
9                   health care from the Administration.

10                  “(G) DEPARTMENT OF DEFENSE AND  
11                  TRICARE PROGRAM.—For purposes of subpara-  
12                  graph (A), the Secretary of Defense shall report  
13                  the sum of—

14                         “(i) the total amount paid for each  
15                         applicable antimicrobial drug procured by  
16                         the Department of Defense for individuals  
17                         who receive health care from the Depart-  
18                         ment; and

19                         “(ii) for each applicable antimicrobial  
20                         drug dispensed under the TRICARE retail  
21                         pharmacy program under section  
22                         1074g(a)(2)(E)(ii) of title 10, United  
23                         States Code, the product of—

24                                 “(I) the per-unit ingredient cost,  
25                                 minus any per-unit rebate paid by the

1 sponsor of the applicable antimicrobial  
2 drug; and

3 “(II) the number of units of such  
4 applicable antimicrobial drug dis-  
5 pensed under such program.

6 “(H) DEPARTMENT OF HOMELAND SECU-  
7 RITY.—For purposes of subparagraph (A), the  
8 Secretary of Homeland Security shall report the  
9 total amount paid for each applicable anti-  
10 microbial drug procured by the Department of  
11 Homeland Security for individuals who receive  
12 health care through a program carried out by  
13 the Department.

14 “(I) BUREAU OF PRISONS.—For purposes  
15 of subparagraph (A), the Director of the Bu-  
16 reau of Prisons shall report the total amount  
17 paid for each applicable antimicrobial drug pro-  
18 cured by the Bureau of Prisons for individuals  
19 who receive health care through the Bureau.

20 “(J) INDIAN HEALTH SERVICE.—For pur-  
21 poses of subparagraph (A), the Secretary, act-  
22 ing through the Indian Health Service, shall re-  
23 port the total amount paid for each applicable  
24 antimicrobial drug procured by the Service for

1 individuals who receive health care through the  
2 Service.

3 “(2) REGULATIONS.—Not later than 1 year  
4 after the date of enactment of this part, the Sec-  
5 retary, in consultation with the heads of Federal  
6 agencies carrying out specified government pro-  
7 grams, shall issue regulations to assist such heads  
8 (or their designees) in carrying out the requirements  
9 under this section.

10 “(3) SUBSCRIPTION CONTRACT ADJUSTMENT.—  
11 Pursuant to the contract entered into under this sec-  
12 tion with respect to an applicable antimicrobial drug,  
13 for each year of the term of such contract, the Sec-  
14 retary shall, not earlier than 6 months after the end  
15 of each calendar year, subtract from the payment in-  
16 stallments determined for such contract under sub-  
17 section (c)(1) for such year the revenue of the spon-  
18 sor of such drug from the previous year from sales  
19 of the applicable antimicrobial drug reported under  
20 paragraph (1) for specified government programs.

21 “(4) DEFINITIONS.—In this subsection:

22 “(A) APPLICABLE ANTIMICROBIAL  
23 DRUG.—The term ‘applicable antimicrobial  
24 drug’ means an antimicrobial drug for which

1 the sponsor of such drug receives a subscription  
2 contract under subsection (a).

3 “(B) SPECIFIED GOVERNMENT PRO-  
4 GRAM.—The term ‘specified government pro-  
5 gram’ means—

6 “(i) the Medicare part D program  
7 under part D of title XVIII of the Social  
8 Security Act;

9 “(ii) the Medicare Part B program  
10 under part B of such title XVIII;

11 “(iii) the Medicare Part A program  
12 under part A of such title XVIII;

13 “(iv) the Medicaid program estab-  
14 lished under title XIX of the Social Secu-  
15 rity Act and includes, with respect to a  
16 State, any waiver in effect with respect to  
17 such program;

18 “(v) any program under which pre-  
19 scription drugs are procured by the De-  
20 partment of Veterans Affairs;

21 “(vi) any program under which pre-  
22 scription drugs are procured by the De-  
23 partment of Defense;



1           “(vii) the TRICARE retail pharmacy  
2           program under section 1074g(a)(2)(E)(ii)  
3           of title 10, United States Code;

4           “(viii) any program under which pre-  
5           scription drugs are procured by the De-  
6           partment of Homeland Security;

7           “(ix) any program under which pre-  
8           scription drugs are procured by the Bu-  
9           reau of Prisons; or

10           “(x) any program under which pre-  
11           scription drugs are procured by the Indian  
12           Health Service.

13           “(e) FAILURE TO ADHERE TO TERMS.—The Sec-  
14           retary shall cease any payment installments under a con-  
15           tract under this section if—

16           “(1) the sponsor—

17           “(A) permanently withdraws the anti-  
18           microbial drug from the market in the United  
19           States;

20           “(B) fails to meet criteria under subsection  
21           (b); or

22           “(C) does not complete a postmarket study  
23           required by the Food and Drug Administration  
24           during the length of the term of the contract;



1 the Director of the Centers for Disease Control and  
2 Prevention shall coordinate with the Administrator  
3 of the Health Resources and Services Administra-  
4 tion, the Administrator of the Centers for Medicare  
5 & Medicaid Services, the National Coordinator for  
6 Health Information Technology, and other relevant  
7 agencies, to establish a grant program under the  
8 Centers for Disease Control and Prevention to sup-  
9 port hospital and other inpatient facility efforts—

10 “(A) to judiciously use antimicrobial drugs,  
11 such as by establishing or implementing appro-  
12 priate use programs, including infectious dis-  
13 ease telehealth programs, using appropriate di-  
14 agnostic tools, partnering with academic hos-  
15 pitals, increasing health care-associated infec-  
16 tion reporting, and monitoring antimicrobial re-  
17 sistance; and

18 “(B) to participate in the National  
19 Healthcare Safety Network Antimicrobial Use  
20 and Resistance Module or the Emerging Infec-  
21 tions Program Healthcare-Associated Infections  
22 Community Interface activity of the Centers for  
23 Disease Control and Prevention or a similar re-  
24 porting program, as specified by the Secretary,  
25 relating to antimicrobial drugs.

1           “(2) PRIORITIZATION.—In awarding grants  
2           under paragraph (1), the Secretary shall prioritize  
3           hospitals without an existing program to judiciously  
4           use antimicrobial drugs, subsection (d) hospitals (as  
5           defined in subparagraph (B) of section 1886(d)(2)  
6           of the Social Security Act that are located in rural  
7           areas (as defined in subparagraph (D) of such sec-  
8           tion), critical access hospitals (as defined in section  
9           1861(mm)(1) of such Act), hospitals serving Tribal-  
10          populations, and safety-net hospitals.

11          “(3) FUNDING.—Of the amounts appropriated  
12          under section 399SS, the Secretary shall reserve  
13          \$500,000,000 to carry out this subsection.

14          “(b) SURVEILLANCE AND REPORTING OF ANTIBIOTIC  
15          USE AND RESISTANCE.—

16          “(1) IN GENERAL.—The Secretary, acting  
17          through the Director of the Centers for Disease  
18          Control and Prevention, shall use the National  
19          Healthcare Safety Network and other appropriate  
20          surveillance systems to assess—

21                  “(A) appropriate conditions, outcomes, and  
22                  measures causally related to antibacterial resist-  
23                  ance, including types of infections, the causes  
24                  for infections, and whether infections are ac-  
25                  quired in a community or hospital setting, in-

1           creased lengths of hospital stay, increased costs,  
2           and rates of mortality; and

3           “(B) changes in bacterial resistance to  
4           antimicrobial drugs in relation to patient out-  
5           comes, including changes in percent resistance,  
6           prevalence of antibiotic-resistant infections, and  
7           other such changes.

8           “(2) ANTIBIOTIC USE DATA.—The Secretary,  
9           acting through the Director of the Centers for Dis-  
10          ease Control and Prevention, shall work with Fed-  
11          eral agencies (including the Department of Veterans  
12          Affairs, the Department of Defense, the Department  
13          of Homeland Security, the Bureau of Prisons, the  
14          Indian Health Service, and the Centers for Medicare  
15          & Medicaid Services), private vendors, health care  
16          organizations, pharmacy benefit managers, and  
17          other entities as appropriate to obtain reliable and  
18          comparable human antibiotic drug consumption data  
19          (including, as available and appropriate, volume an-  
20          tibiotic distribution data and antibiotic use data, in-  
21          cluding prescription data) by State or metropolitan  
22          areas.

23          “(3) ANTIBIOTIC RESISTANCE TREND DATA.—  
24          The Secretary, acting through the Director of the  
25          Centers for Disease Control and Prevention, shall in-

1       tensify and expand efforts to collect antibiotic resist-  
2       ance data and encourage adoption of the Antibiotic  
3       Use and Resistance Module within the National  
4       Healthcare Safety Network among all health care fa-  
5       cilities across the continuum of care, including, as  
6       appropriate, acute care hospitals, dialysis facilities,  
7       nursing homes, ambulatory surgical centers, and  
8       other ambulatory health care settings in which anti-  
9       microbial drugs are routinely prescribed. The Sec-  
10      retary shall seek to collect such data from electronic  
11      medication administration reports and laboratory  
12      systems to produce the reports described in para-  
13      graph (4).

14           “(4) PUBLIC AVAILABILITY OF DATA.—The  
15      Secretary, acting through the Director of the Cen-  
16      ters for Disease Control and Prevention, shall, for  
17      the purposes of improving the monitoring of impor-  
18      tant trends in patient outcomes in relation to anti-  
19      bacterial resistance—

20           “(A) make the data derived from surveil-  
21      lance under this subsection publicly available  
22      through reports issued on a regular basis that  
23      is not less than annually; and

24           “(B) examine opportunities to make such  
25      data available in near real time.

1 **“SEC. 399SS. APPROPRIATIONS.**

2 “(a) IN GENERAL.—To carry out this part, there are  
3 hereby appropriated to the Secretary, out of amounts in  
4 the Treasury not otherwise appropriated,  
5 \$11,000,000,000, for fiscal year 2022, to remain available  
6 until expended.

7 “(b) EMERGENCY DESIGNATION.—

8 “(1) IN GENERAL.—The amounts provided by  
9 this section are designated as an emergency require-  
10 ment pursuant to section 4(g) of the Statutory Pay-  
11 As-You-Go Act of 2010.

12 “(2) DESIGNATION IN SENATE.—In the Senate,  
13 this section is designated as an emergency require-  
14 ment pursuant to section 4112(a) of H. Con. Res.  
15 71 (115th Congress), the concurrent resolution on  
16 the budget for fiscal year 2018.

17 **“SEC. 399TT. STUDIES AND REPORTS.**

18 “(a) IN GENERAL.—Not later than 6 years after the  
19 date of enactment of this part, the Comptroller General  
20 of the United States shall complete a study on the effec-  
21 tiveness of this part in developing priority antimicrobial  
22 drugs. Such study shall examine the indications for, usage  
23 of, development of resistance with respect to, and private  
24 and societal value of critical need antimicrobial drugs, and  
25 the impact of the programs under this part on patients  
26 and markets of critical need antimicrobial drugs. The

1 Comptroller General shall report to the Committee on  
2 Health, Education, Labor, and Pensions of the Senate and  
3 the Committee on Energy and Commerce of the House  
4 of Representatives on the findings of such study.

5 “(b) ANTIBIOTIC USE IN THE UNITED STATES; AN-  
6 NUAL REPORTS.—The Director of the Centers for Disease  
7 Control and Prevention shall, each year, update the report  
8 entitled ‘Antibiotic Use in the United States’ to include  
9 updated information on progress and opportunities with  
10 respect to data, programs, and resources for prescribers  
11 to promote appropriate use of antimicrobial drugs.

12 “(c) REPORT ON ANTIMICROBIAL PROPHYLACTICS.—  
13 Not later than 3 years after the date of enactment of this  
14 part, the Director of the Centers for Disease Control and  
15 Prevention shall publish a report on antimicrobial prophyl-  
16 lactics.

17 **“SEC. 399UU. DEFINITIONS.**

18 “In this part—

19 “(1) the term ‘antimicrobial drug’—

20 “(A) means, subject to subparagraph (B),  
21 a product that is—

22 “(i) a drug that directly inhibits rep-  
23 lication of or kills bacteria or fungi rel-  
24 evant to the proposed indication at con-  
25 centrations likely to be attainable in hu-



1           mans to achieve the intended therapeutic  
2           effect; or

3                   “(ii) a biological product that acts di-  
4                   rectly on bacteria or fungi or on the sub-  
5                   stances produced by such bacteria or fungi;  
6                   and

7                   “(B) does not include—

8                           “(i) a drug that achieves the effect de-  
9                           scribed by subparagraph (A)(i) only at a  
10                          concentration that cannot reasonably be  
11                          studied in humans because of its antici-  
12                          pated toxicity; or

13                           “(ii) a vaccine; and

14                           “(2) the term ‘Committee’ means the Com-  
15                          mittee on Critical Need Antimicrobials established  
16                          under section 39900.”.

○