### 116TH CONGRESS 1ST SESSION

# H. R. 2620

To advance treatment and cures for blindness and other retinal conditions and to promote competitiveness in the United States through a pilot program to increase funding for translational research, and for other purposes.

### IN THE HOUSE OF REPRESENTATIVES

May 9, 2019

Mr. BISHOP of Georgia (for himself, Mrs. Rodgers of Washington, Mr. BILI-RAKIS, Mr. COHEN, Mr. O'HALLERAN, Mr. SCHNEIDER, and Mr. FITZPATRICK) introduced the following bill; which was referred to the Committee on Energy and Commerce

# A BILL

To advance treatment and cures for blindness and other retinal conditions and to promote competitiveness in the United States through a pilot program to increase funding for translational research, 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 "Faster Treatments and
- 5 Cures for Eye Diseases Act".
- 6 SEC. 2. DEFINITIONS.
- 7 For purposes of this Act:

- 1 (1)APPLICABLE CONGRESSIONAL COMMIT-2 TEES.—The term "applicable congressional commit-3 tees" means the Committees on Energy and Com-4 merce and Financial Services of the House of Rep-5 resentatives and the Committees on Banking, Hous-6 ing, and Urban Affairs and Health, Education, 7 Labor, and Pensions of the Senate.
- 8 (2) APPROPRIATE FEDERAL BANKING AGEN9 CY.—The term "appropriate Federal banking agen10 cy" has the meaning given that term in section 3 of
  11 the Federal Deposit Insurance Act (12 U.S.C.
  12 1813).
  - (3) Cost.—The term "cost" has the meaning given to the term "cost of a loan guarantee" in section 502(5)(C) of the Federal Credit Reform Act of 1990 (2 U.S.C. 661a(5)(C)).
  - (4) Depository institution; depository institution holding company.—The term "depository institution" and "depository institution holding company" have the meaning given those terms under section 3 of the Federal Deposit Insurance Act (12 U.S.C. 1813).
- 23 (5) EYE BOND.—The term "eye bond" means a bond—

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1	(A) issued by an issuer pursuant to this
2	Act;
3	(B) the proceeds of which are used to fund
4	projects selected under section 6(b)(1), except
5	as otherwise described in this Act; and
6	(C) that complies with the regulations
7	issued under section 4.
8	(6) Funded Project.—The term "funded
9	project" means a translational research project that
10	is selected to be funded using the proceeds of an eye
11	bond.
12	(7) Guarantee.—The term "guarantee" has
13	the meaning given to the term "loan guarantee" in
14	section 502 of the Federal Credit Reform Act of
15	1990 (2 U.S.C. 661a) and includes a loan guarantee
16	commitment (as defined in such section 502).
17	(8) Issuer.—The term "issuer" means an enti-
18	ty that—
19	(A) is a depository institution, a depository
20	institution holding company, or a broker or
21	dealer registered with the Securities and Ex-
22	change Commission;
23	(B) complies with the schedule for the
24	issuance of eye bonds established under section
25	3(e); and

1	(C) complies with the regulations issued
2	under section 5.
3	(9) Program.—The term "Program" means
4	the Eye Bond Pilot Program established under sec-
5	tion 3.
6	(10) Secretary.—The term "Secretary"
7	means the Secretary of Health and Human Services,
8	except in references to the Secretary of the Treas-
9	ury.
10	(11) State.—The term "State" means—
11	(A) each State of the United States;
12	(B) the District of Columbia;
13	(C) the Commonwealth of Puerto Rico;
14	and
15	(D) any other territory or possession of the
16	United States.
17	(12) Translational research.—The term
18	"translational research"—
19	(A) means any research project that is de-
20	signed to cure vision blindness and any condi-
21	tions attendant to vision impairment that are,
22	as determined by the Director of the National
23	Eye Institute, congenital to the vision impair-
24	ment and not incidental to vision impairment or
25	caused by vision impairment;

1	(B) includes projects designed to cure—
2	(i) hearing impairment genetically
3	linked to vision impairment, such as Usher
4	Syndrome;
5	(ii) retinal degenerative diseases such
6	as retinitis pigmentosa, macular degenera-
7	tion, and Usher Syndrome;
8	(iii) vision trauma due to injury such
9	as that experienced by wounded veterans;
10	(iv) glaucoma;
11	(v) optic nerve disorders that result in
12	vision impairment or blindness, such as
13	morning glory syndrome; and
14	(vi) diabetic reinopathy; and
15	(C) subject to subparagraph (B)(vi), does
16	not include projects designed to cure any under-
17	lying disease or condition whose symptoms may
18	include vision impairment, such as diabetes.
19	SEC. 3. EYE BOND PILOT PROGRAM.
20	(a) Establishment.—Not earlier than 1 year after
21	the date of enactment of this Act, the Secretary, in con-
22	sultation with the Secretary of the Treasury and the ap-
23	propriate Federal banking agencies, shall establish a pilot
24	program to be known as the Eye Bond Pilot Program
25	under which—

1	(1) the Director of the National Eye Institute
2	shall, in accordance with section 6, select
3	translational research projects to be funded by eye
4	bonds; and
5	(2) the Secretary shall—
6	(A) provide a partial Federal guarantee of
7	the eye bonds;
8	(B) contract with an issuer to issue the eye
9	bonds; and
10	(C) use the proceeds from the sale of the
11	eye bonds to fund the selected projects and to
12	pay for other related expenses, as permitted by
13	this Act.
14	(b) Federal Guarantee.—
15	(1) In General.—The Secretary shall guar-
16	antee the payment of principal (but not the payment
17	of interest) on an eye bond, on a bond-by-bond
18	basis, in an amount to be determined by the Sec-
19	retary, but in no case may the amount of such guar-
20	antee be more than 50 percent of the principal of
21	the eye bond.
22	(2) Prioritization of Taxpayer inter-
23	ESTS.—All eye bonds shall be structured to give first
24	priority to protecting the interests of the United

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States by ensuring that—

1	(A) all cash proceeds received from the re-
2	payment of an eye bond are first used to reduce
3	the amount of principal guaranteed by the Sec-
4	retary under the terms of the eye bond; and
5	(B) the Secretary has a senior claim on all
6	assets and collateral under the eye bond to the
7	extent the guarantee provided by the Secretary
8	is not extinguished.
9	(3) Limitation.—The Secretary may not guar-
10	antee—
11	(A) more than \$1,000,000,000 for all eye
12	bonds issued pursuant to this Act; and
13	(B) more than \$250,000,000 for all eye
14	bonds issued pursuant to this Act in any single
15	fiscal year.
16	(c) Issuance Schedule.—The Secretary shall, in
17	consultation with the Secretary of the Treasury and
18	issuers, establish a schedule for the issuance of eye bonds
19	that ensures that funded projects represent a reasonable
20	sample of diverse causes of vision loss.
21	(d) Risk-Share Pool.—
22	(1) In general.—With respect to an eye bond
23	guaranteed under this section, the Secretary may
24	allow a risk-share pool capitalized by issuers to pro-
25	vide a first-loss guarantee of the principal and inter-

1	est of such bond, if the Secretary determines that
2	such first-loss guarantee would—
3	(A) be a robust source of protection for the
4	United States, as guarantor of the eye bond,
5	that reduces the cost of the Federal guarantee
6	to the United States;
7	(B) encourage the flow of private sector
8	capital into biomedical translational research;
9	(C) create a prudent incentive for issuers
10	to contribute additional private capital for bio-
11	medical translational research; or
12	(D) meet other public interest, safety, and
13	soundness goals.
14	(2) Consultation.—In making a determina-
15	tion under paragraph (1), the Secretary shall consult
16	with the Secretary of the Treasury and the appro-
17	priate Federal banking agencies.
18	(3) Funding.—The cost of contracting with a
19	risk-share pool under this subsection shall be paid
20	from the proceeds from the sale of eye bonds pursu-
21	ant to the Program.
22	(e) Equity Position Option for the Sec-
23	RETARY.—
24	(1) In general.—With respect to an eye bond
25	issued pursuant to this Act, the Secretary, in con-

sultation with the Secretary of the Treasury and other appropriate parties, including eye bond issuers and investors, may negotiate an equity position for the United States Government in the projects to be funded by such eye bond if the Secretary determines that such an equity position will further the interests of the Program and the United States.

## (2) Limitations.—

- (A) SUPPLEMENTAL.—Any equity position taken in a project pursuant to paragraph (1) shall be supplemental to, not in lieu of, a guarantee provided by the Secretary with respect to the eye bond funding such project.
- (B) Total amount.—The total of amount of an eye bond guarantee under this section and any equity position taken by the Secretary in a project funded by such bond that is supplemental to such guarantee shall not exceed 50 percent of the principal amount of the eye bond.
- (3) Notification and consultation.—Prior to finalization of any equity position under paragraph (1), the Secretary shall notify the applicable congressional committees and consult with such committees on the proposed terms of such equity po-

1	sition and whether taking such equity position will
2	further the interests of the Program and the United
3	States.
4	(f) TERMINATION OF THE PROGRAM.—
5	(1) In general.—Except as provided in para-
6	graph (2), the Program shall terminate on the date
7	that is 6 years after the date of enactment of this
8	Act.
9	(2) Early termination.—
10	(A) IN GENERAL.—The Secretary may ter-
11	minate the Program before the date described
12	in paragraph (1).
13	(B) Congressional notification.—If
14	the Secretary determines that the Program
15	shall be terminated under subparagraph (A),
16	not later than 60 days before the date on which
17	the termination is effective, the Secretary
18	shall—
19	(i) submit to the applicable congres-
20	sional committees a report that includes—
21	(I) a description of the reasons
22	for the termination;
23	(II) any corrective actions that
24	may be taken; and

1	(III) any other actions that may
2	be taken to promote the use of private
3	capital and to increase the amount of
4	Federal funds made available to carry
5	out basic and translational biomedical
6	research; and
7	(ii) make publicly available the report
8	described in clause (i).
9	(3) Effect on eye bonds issued and fed-
10	ERAL GUARANTEES.—The termination of the Pro-
11	gram shall not affect the validity of—
12	(A) any eye bond issued before the date on
13	which the Program is terminated; or
14	(B) any Federal guarantee under this Act
15	for an eye bond described in subparagraph (A).
16	(g) Program Funding.—
17	(1) Administrative expenses paid from
18	BOND SALES.—Except as provided under paragraph
19	(2), the cost of carrying out this Act, including the
20	cost to the Secretary in administering the Program,
21	shall be recovered from the proceeds from the sale
22	of eye bonds pursuant to the Program or from fees
23	as set forth in paragraph (3).

1	(2) Specific appropriation or contribu-
2	TION.—No guarantee shall be made under this sec-
3	tion unless—
4	(A) an appropriation for the full cost of
5	the guarantee has been made;
6	(B) the Secretary has received from the
7	eye bond issuer a payment in full for the cost
8	of the guarantee; or
9	(C) a combination of an appropriation and
10	the deposit of a payment from the bond issuer
11	into the Treasury has been made in a sufficient
12	amount to cover the full cost of the guarantee.
13	(3) Cost of guarantees.—
14	(A) IN GENERAL.—The Secretary shall
15	charge and collect fees for guarantees under
16	this section in amounts the Secretary deter-
17	mines are sufficient to recover applicable ad-
18	ministrative expenses.
19	(B) AVAILABILITY.—Fees collected under
20	this subsection—
21	(i) shall be deposited by the Secretary
22	into the Treasury; and
23	(ii) are authorized to remain available
24	until expended.

# SEC. 4. EYE BOND TERMS, CONDITIONS, AND STRUCTURE.

- 2 (a) IN GENERAL.—Not later than 180 days after the
- 3 date of enactment of this Act, the Secretary, in consulta-
- 4 tion with the Secretary of the Treasury, the Chairman of
- 5 the Securities and Exchange Commission, the heads of the
- 6 appropriate Federal banking agencies, the Director of the
- 7 National Institutes of Health, and the heads of other Fed-
- 8 eral departments and agencies and other interested parties
- 9 as the Secretary determines appropriate, shall issue regu-
- 10 lations to specify the terms, conditions, and structure for
- 11 an eye bond.
- 12 (b) Auctions.—In issuing the regulations under
- 13 subsection (a)—
- 14 (1) the Secretary may provide for an auction to
- select the purchasers of eye bonds; and
- 16 (2) any such auction may include a process that
- minimizes the risk to the Government of the Federal
- guarantee involved by allowing bidders for an eye
- bond to compete against each other by bidding on
- the percentage of the Federal guarantee under sec-
- 21 tion 4(b) with respect to the eye bond, with the bid
- for the lowest percentage winning the auction, tak-
- 23 ing into account other terms and conditions set by
- 24 the issuer to ensure the lowest total cost to the Gov-
- ernment.

# 1 SEC. 5. EYE BOND ISSUERS.

2	(a) Issuer Criteria.—Not later than 180 days
3	after the date of enactment of this Act, the Secretary, in
4	consultation with the Secretary of the Treasury, shall
5	issue regulations—
6	(1) to establish the criteria for selecting issuers;
7	(2) to ensure that issuers perform in a manner
8	that ensures the successful issuance of eye bonds
9	that promote biomedical translational research in the
10	United States; and
11	(3) to ensure that issuers use sound under-
12	writing practices that protect the interests of—
13	(A) the United States;
14	(B) eye bond investors; and
15	(C) the long-term promotion of
16	translational research for vision impairment
17	and other diseases, disabilities, and syndromes
18	congenital to vision impairment or caused by vi-
19	sion impairment, taking into account features
20	that are valuable after any authorization for ex-
21	panded use of a limited Federal guarantee for
22	biomedical translational research for other dis-
23	eases and disabilities.
24	(b) Compensation for Issuers.—The issuer of an
25	eye bond shall be compensated from the proceeds from the

1	sale of such eye bond at such rate and on such terms as
2	the Secretary may provide.
3	(c) Public Disclosures With Respect to Eye
4	Bonds.—
5	(1) In general.—Not less than 2 business
6	days before the date on which an issuer issues an
7	eye bond, the issuer shall file with the Securities and
8	Exchange Commission, and make available to the
9	public, the following information:
10	(A) The nature of all projects funded by
11	the eye bond.
12	(B) The name of any principal individual
13	or institution that will be conducting each
14	project.
15	(C) The milestones established for each
16	project.
17	(D) A determination by the issuer as to
18	whether each project funded by the eye bond
19	has appropriately protected intellectual prop-
20	erty.
21	(E) The structure of the eye bond.
22	(F) The interest payment schedule for the
23	eye bond.
24	(G) The anticipated returns and risks of
25	the eye bond.

1 (H) Such other information as the Com-2 mission determines necessary or appropriate in 3 the public interest or for the protection of in-4 vestors.

#### (2) Rulemaking.—

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- (A) IN GENERAL.—Not later than 180 days after the date of enactment of this Act, the Securities and Exchange Commission shall issue regulations to carry out this subsection.
- (B) AUTHORITY OF THE SECRETARY.—If the Securities and Exchange Commission does not issue the regulations required under subparagraph (A) before the end of the 180-day period described under subparagraph (A), the Secretary shall issue regulations to carry out this subsection before the end of the 60-day period beginning on the end of the 180-day period described under subparagraph (A).

#### 19 SEC. 6. TRANSLATIONAL RESEARCH PROJECTS.

20 (a) ELIGIBILITY REQUIREMENTS.—Not later than
21 180 days after the date of enactment of this Act, the Sec22 retary, in consultation with the Secretary of the Treasury,
23 the Director of the National Eye Institute, and other in24 terested parties, shall issue final regulations for the eligi25 bility criteria for selecting translational research projects

1	that will be funded through eye bonds. Such regulations
2	shall address—
3	(1) the stage of clinical trial for projects to pro-
4	vide the greatest likelihood of commercial applica-
5	tion;
6	(2) the variations among disease and conditions
7	needed to ensure sufficient diversification in each
8	eye bond; and
9	(3) the number of possible cures and treat-
10	ments that are needed as determined by the Sec-
11	retary, in consultation with issuers and the Director
12	of the National Eye Institute, to ensure the success-
13	ful issuance of eye bonds so as to protect the United
14	States as guarantor of the eye bonds, including—
15	(A) drug therapies;
16	(B) gene therapies; and
17	(C) artificial restoration of sight and simi-
18	lar mechanisms.
19	(b) Selection of Projects.—
20	(1) In general.—The Director of the National
21	Eye Institute, in consultation with the Director of
22	the National Institutes of Health and the Secretary
23	of the Treasury, shall select translational research
24	projects to be funded with the proceeds of an eye
25	bond.

1	(2) Factors for selection.—Not later than
2	30 days after the date on which the final regulations
3	are issued under subsection (a), the Secretary shall
4	submit to the Director of the National Eye Institute
5	factors that the Director of the National Eye Insti-
6	tute shall consider in making the selection under
7	paragraph (1), including—
8	(A) the amount of equity any intellectual
9	property holder will hold in the project;
10	(B) the resources any individual or institu-
11	tion will be required to demonstrate to ensure
12	the ability of the individual or institution to
13	repay the obligation under the eye bond, re-
14	gardless of the success or failure of the project
15	funded with the proceeds of the eye bond;
16	(C) the number of projects needed to en-
17	sure diversification of risk;
18	(D) the manner in which funded projects
19	will be defunded if the interim goals of the
20	project are not satisfied; and
21	(E) such other factors related to bio-
22	medical translational research project selection
23	as the Secretary determines appropriate.
24	(3) Additional consultations.—

- 1 (A) IN GENERAL.—In carrying out para2 graph (1), the Director of the National Eye In3 stitute may establish any consultative body that
  4 the Director determines is necessary to provide
  5 for a complete, transparent, and forward-look6 ing selection of projects.
  - (B) Scientific advisers.—In carrying out paragraph (1), the Director of the National Eye Institute may consult with any group of scientific advisers that the Director determines is necessary.

## (c) Establishment of Milestones.—

- (1) IN GENERAL.—The Director of the National Eye Institute shall, for each project funded by an eye bond, establish milestones to determine the probability of success or failure for such project.
- (2) Inclusion in filings.—The Director of the National Eye Institute shall submit to the issuer of an eye bond the milestones for each project funded from such eye bond, so such milestones may be included in the filings made available by the issuer to the public under section 5(c).
- 23 (d) RESEARCH REQUIREMENT.—Translational re-24 search carried out under a project funded by an eye bond 25 shall be conducted—

1	(1) in a State; and
2	(2) by an individual or institution that is—
3	(A) chartered in accordance with the laws
4	of that State; and
5	(B) clearly subject to verification of bene-
6	ficial ownership by the issuer and, upon re-
7	quest, by the Secretary.
8	SEC. 7. INAPPLICABILITY OF CERTAIN LAWS.
9	Eye bonds shall not be subject to—
10	(1) section 15G of the Securities Exchange Act
11	of 1934 (15 U.S.C. 780–11);
12	(2) except as provided under section 5(c), any
13	registration or disclosure requirement promulgated
14	by the Securities and Exchange Commission; and
15	(3) section 13 of the Bank Holding Company
16	Act of 1956 (12 U.S.C. 1851).
17	SEC. 8. REPORTS.
18	(a) GAO STUDY AND REPORTS ON OTHER RE-
19	SEARCH PROJECTS.—
20	(1) Ongoing study.—The Comptroller Gen-
21	eral of the United States shall carry out an ongoing
22	study to consider whether a program similar to the
23	Eye Bond Pilot Program under this Act should be
24	established for other biomedical research projects.

1 (2) Report.—The Comptroller General shall, 2 during the period beginning on the date of the estab-3 lishment of the Program and ending on the termination date of the Program, issue a report to the ap-5 plicable congressional committees, not less frequently 6 than annually, on all findings and determinations 7 made in carrying out the study required under para-8 graph(1). 9 (b) Reports on the Program.—Not later than 2 10 years after the date on which eye bonds are first issued under this Act, and annually thereafter during the period 12 ending on the date that is 4 years after the date on which 13 eye bonds are first issued under this Act, the Comptroller 14 General and the Director of the National Institutes of 15 Health (in consultation with the Director of the National Center for Advancing Translational Sciences) shall each 16 17 issue a separate report to the applicable congressional 18 committees on— 19 (1) the progress of the issuance of eye bonds; 20 (2) the reasons for any problems achieving de-21 sired volumes of eye bonds or the ability of the Pro-22 gram to proceed at a faster pace; 23 (3) an analysis of the risk to the Government 24 in providing the Federal guarantee described under

section 4(b);

1	(4) any improvements to eye bonds that the
2	Secretary should consider;

- (5) the applicability of financial instruments similar to eye bonds to other biomedical research areas such as cancer, Alzheimer's disease, rare diseases or conditions (as defined in section 526 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360bb)) and syndromes of particular concern to children; and
- (6) any other matter that the Comptroller General or the Director, respectively, determines is appropriate.

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