The Foundation for the National Institutes of Health, a non-profit, 501(c)(3) charitable organization that supports the National Institutes of Health (NIH) in its mission to improve health by forming and facilitating public-private partnerships for biomedical research, is issuing a Request for Proposals (RFP) to support the Accelerating Medicines Partnership® (AMP®) Bespoke Gene Therapy Consortium (BGTC). The BGTC is executed by FNIH as a public-private partnership involving the NIH, the U.S. Food and Drug Administration (FDA), multiple pharmaceutical companies, non-profits and patient advocates and is dedicated to making gene therapy a reality for people with rare genetic diseases affecting populations too small to be viable from the current commercial perspective.

Despite the potential for substantial therapeutic impact, the limited access to bespoke, or tailor-made, gene therapy for such diseases has been recognized by pharmaceutical, academic, NIH and FDA leadership. Ongoing discussions to address this issue have focused on using well-established adeno-associated virus (AAV) vectors as the single delivery technology, and the ability to facilitate and generalize access and processes for the development of bespoke gene therapies. In order to accelerate and advance the scalability, reproducibility, and regulatory ability to deliver individualized, or “bespoke” gene therapies to ultra-rare patient populations, the BGTC will combine studies focused on AAV biology as related to human gene therapy with a pilot clinical trial program manufacturing and testing AAV vectors in a more standardized fashion. The BGTC expects to use the research and clinical data to produce an operational playbook that invokes the use of streamlined templates, master regulatory files, and uniform production processes to inform and streamline future gene therapies. This model program provides a tremendous opportunity to impact patients with very rare diseases and offers hope for a pathway toward the commercial viability of these treatments.
Purpose

The overall goal for this RFP is to solicit proposals for assay development and other approaches that will advance understanding of the mechanistic steps between transduction of a target cell population with an AAV vector and accumulation of the therapeutic gene product in target cells, with the ultimate goal of enhancing the therapeutic impact of AAV gene therapy in humans. The primary focus of this RFP is for the development of high-throughput screening (HTS)-compatible assays for these individual mechanistic steps, which could be used to screen small molecule libraries to identify optimization strategies for individual mechanistic steps in AAV transduction and targeted gene expression. However, proposals for alternate, high-impact approaches to enhance the therapeutic efficacy of AAV gene therapy in humans will be considered.

Background

Advancing the understanding of AAV biology as related to human gene therapy has been identified as an important pre-competitive area that could benefit from a focused public-private partnership. Many areas of AAV biology, such as viral genome packaging, virion assembly, virion release, cell fusion, intracellular transport, and the regulation of therapeutic gene expression, remain incompletely understood. It is thought that the efficiency in manufacturing of AAV therapies, as well as therapeutic efficacy in patients, could be substantially improved by a better understanding of AAV biology, and optimization of key mechanistic steps. This RFP (2021-BGTC002) is focused on developing methods to interrogate the biology of AAV in target cells to enhance therapeutic gene expression in patients. Results will ultimately be used to identify gene expression improvements in target cells related to diseases/disorders selected for the BGTC clinical program. A companion RFP (2021-BGTC001) will focus on AAV biology in host producer cells to increase the efficiency of producing high-quality clinical grade AAV gene therapy vectors.

Specific Research Objectives

This RFP solicits proposals to substantially increase the expression of a therapeutic gene delivered by an AAV vector in target cells. The overarching technical objective for responses to this RFP within the scientific topic areas described below is to develop solutions which improve one or more steps in the recombinant AAV gene expression process. Standardization of the proposed solution or assay is critical, and the responses should outline the planned studies to be conducted to address this requirement. Submissions proposing the development of HTS-compatible assays must be applicable to any human cell type and must be designed such that assay(s) developed could be incorporated into existing NCATS screening capabilities by the conclusion of the term of the award. For further details, see Assay Development & Screening | National Center for Advancing Translational Sciences (nih.gov).

The topic area of investigation targeted by this RFP is improvement of therapeutic gene expression from recombinant AAV vectors in human gene therapy, which may include (but is not limited to) assays and other methods for research on and optimization of the following mechanistic steps: internalization of the vector, the endosomal state of the AAV vector; trafficking to the nucleus; uncoating in the nucleus; second strand synthesis; concatemerization of the viral genome; regulation of therapeutic gene expression events, appropriate post-translational modification and trafficking of gene products. HTS-compatible assays for one or more of the mechanistic steps described above are of particular interest. Proposals may also be focused more broadly on other approaches for increasing the efficacy of AAV gene therapy in humans. Research focused on the host immune response would be considered out of scope for this initiative (please see the Accelerating Research and Development for Advanced Therapies (ARDAT) project being conducted by the Innovative Medicines Initiative for more information on research efforts in this space).
In addition to overall scientific merit and rigor, review criteria will include the following considerations:

- Is the work proposed focused on the mechanistic steps between entry of AAV in a target cell and between therapeutic gene expression and processing/trafficking and accumulation of gene products?
- Is the work likely to result in a clinically relevant enhancement of therapeutic gene expression in patients?
- For HTS assays, are they compatible with the NCATS screening platform?
- Is the proposed work likely to result in communicable deliverables within the project period?

Additional funding considerations include:

- Availability of funds
- Current state of knowledge in the field at the time of funding
- Overall BGTC program balance and lack of overlap with other projects

**Award Information**

I. **Funds Available and Anticipated Number of Awards**

The number of awards and the amount per award is contingent upon the submission of a sufficient number of meritorious submissions and proper budget justification within the proposal.

II. **Award Budget**

Proposal budgets are limited to an amount up to $250,000 of direct costs per year for up to two (2) years and need to reflect the actual needs of the development of the proposed assay. Indirect costs (F&A) must be 15% or less. Proper scientific and budget justification will need to be provided for evaluation. The proposal review working group reserves the right to award at a lower amount than requested.

III. **Award Project Period**

The scope of the proposed assay work should determine the award project period. The request may be for up to two (2) years of funding. The earliest anticipated start date is April 1, 2022.

**Eligibility Information**

Organizations eligible to apply are:

- Private or public sector
- US-based or international
- Able to comply with the necessary AMP BGTC intellectual property, data sharing, and publication guidelines (AMP BGTC Principles can be found in Appendix 2).
Proposal and Submission Instructions

I. Submission Deliverables

Complete proposals will include:

- Proposal research plan, which should describe the information below, but more details can be provided in the proposal response template (Appendix 1):
  - Background, specific aims, preliminary studies (if applicable), research methods, other relevant supporting documents (optional)
  - For proposals other than development of HTS-compatible screening assays: Justification of the broader applicability of the targeted biology and rationale for why this assay would benefit the AAV field beyond the specified disorder
  - For HTS-compatible screening assays, proposals must address compatibility with NCATS HTS-assay guidance criteria (see: https://ncats.nih.gov/preclinical/drugdev/assay#criteria).
- Detailed budget that delineates:
  - Personnel
  - Reagents, materials, equipment, sample acquisition (if necessary)
  - Other requirements for work proposed
  An example budget table can be found in Appendix 1
- Detailed budget justification
- Proposed project timeline
- Biosketches for the Principal Investigators.

II. Data, Publications and Intellectual Property

All applicants will be expected to comply with the AMP BGTC Principles that have already been established for the partnership. These are available in Appendix 2 and will be attached to any award agreements for those projects selected for funding.

III. Page Limit

Please keep your Research plan responses under 10 pages in length (single spaced, 11-point Arial or Times New Roman font), including figures and legends.
Biosketches for the Principal Investigators should not exceed 3 pages.
Further section length requirements are provided in Appendix 1.

IV. Award Reporting

For those proposals selected for award, the Principal Investigators on the award should expect to submit progress updates for the project every 6 months in a format that will be described in the award agreements.

V. Additional Information Required

Please provide any existing IP or patent information relevant to the assay that may affect its use in the partnership, or the banking of any resulting data funded by this effort in a public controlled access database for use after initial publication of the findings. Further guidance is available upon request.
VI. Submission Instructions

Send responses via e-mail to BGTC@fnih.org with “2021-BGTC002 AAV Therapeutic Gene Expression Proposal” in the subject line.

Key Dates

Submission Due Date: February 18, 2022, 11:59 PM Eastern Time


Potential Oral Presentations from Finalists (If Needed): Date TBD
Applicants will be informed after initial review of proposals whether they will need to provide an oral presentation with the ability for Q&A to the BGTC Proposal Review Working Group

Targeted Award Announcement: April 2022
Applicants will be notified by email of the outcome of the RFP.

About the Foundation for the NIH
Established by the United States Congress to support the mission of the NIH – improving health through scientific discovery in the search for cure – the Foundation for the NIH is a leader in identifying and addressing complex scientific and health issues. The Foundation is a non-profit, 501(c)(3) charitable organization that raises private-sector funds for and manages a broad portfolio of unique programs that complement and enhance NIH priorities and activities. For additional information about the Foundation for the NIH, visit www.fnih.org.
Appendix 1

BGTC Recombinant AAV Gene Expression RFP Submission Form

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<th>Title of Project</th>
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<tr>
<td><strong>Principal Investigator(s)</strong> (submitter)</td>
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<td>Name:</td>
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<td>Submitting Organization:</td>
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| Co-investigator(s) (add more as needed) |
| Name: |
| Title: |
| Submitting Organization: |
| Address: |
| e-mail: |
| Tel: |

| Submission Date: |
| Time Period of Project: |
| Project Total Budget: |

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<tr>
<th>Internal Use Only</th>
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<tbody>
<tr>
<td>WG Decision, Date:</td>
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<tr>
<td>Steering Committee Decision, Date: <em>(if needed)</em></td>
</tr>
<tr>
<td>Executive Committee Decision, Date: <em>(if needed)</em></td>
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</tbody>
</table>
Section 1: Project Overview (500 words maximum)
Describe the assay to be developed, the clinical/scientific need for the assay, and the capability gap being addressed.

1.1 Specific Aims and Objectives

1.2 Project Deliverables/Outputs

1.3 Applicability - Describe why this project is appropriate for the BGTC RFP, the pre-competitive nature of the project, and how the project is novel.
- Include a brief description of other efforts known to the applicant to ensure non-duplication.
- Does the work proposed have scientific merit?
- Is the work proposed focused on developing assays or other methods to interrogate the biology of AAV capsids and cargo in the events that follow receptor binding on the surface of a target cell to enhance therapeutic gene expression in a patient?
- Is the work likely to stimulate additional activity leading to progress in understanding the life cycle of AAV? Other?
- Will the proposed work lead to communicable deliverables within the project period?
- Provide an estimate and calculation for the % improvement that can be expected in efficacy or other metric, as a result of this work.

Section 2: Scientific Design (2000 words maximum)

2.1 Background and Supporting Data
- Including Precision, reproducibility, target tissue

2.2 Experimental Plan
- Describe in detail the scientific strategy, design and logistics. Include how the study design will address the project goals and objectives.
- Describe in detail the experiments to be conducted
- Describe the type of data to be analyzed (retrospective and/or prospective)

2.3 Analytical Methods
- Describe in detail analytical methods that will be used.
- Provide a statistical analysis plan, including power calculations.
- If any studies are designed to replicate preliminary data or findings from prior studies, please describe those clearly.

2.4 Technologies and Assays
- Describe the current technologies or assays to be used in the project.
- If project is in Stage 3 or Stage 4, please provide analytical validation data for the assay(s) to be used.

2.5 Human/Animal Subjects
- Will human or animal subjects be involved in this Project? If so, how?
Section 3: Data Sharing and Intellectual Property Management Plan (1000 words maximum)

Please note: Use of pre-existing IP and sharing of newly generated IP must comply with AMP BGTC Principles in Appendix 2.

3.1 Describe pre-existing intellectual property (IP) that could have a bearing on the project.

3.2 List any relevant existing patents and patent applications held by key participants in the project (include patent number, title, submission date).

3.3 Describe any new IP that may be generated.

3.4 Describe plans for risk management concerning:
   - Data use
   - Data security
   - Legal compliance

3.5 Describe plans for depositing data in the BGTC Data Portal or a similarly approved public database prior to initial publication.

Please note: Data sharing plans must comply with AMP BGTC Principles in Appendix 2.

Section 4: Timeline, Milestones, Deliverables and Budget (1000 words maximum)

4.1 Provide a timeline for deliverables, and an end date for the project.

4.2 Describe the milestones and how will progress on achieving them will be assessed.
   - Describe any opportunities for interim feasibility assessment(s) based on Project progress.

4.3 Project Budget
   - Please complete the high-level budget table (see next page) in addition to attaching a detailed budget justification explaining the need for the costs requested with detailed breakdown of costs. Submission will not be reviewed if both of these items are not included.
<table>
<thead>
<tr>
<th>Category</th>
<th>% FTE</th>
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4.4 Detail any existing funding relationships.

Section 5: Program Support *(500 words maximum)*

5.1 List key personnel, including names, titles and role in the Project.
   • List collaborators, advisors, and/or hired consultants outside the consortium. Include letters of support.

Section 6: Legal Agreements

6.1 Legal Agreements
   • List any contracts, memoranda of understanding, grants, data or material transfer agreements that are likely to be necessary to execute the project plan.

Section 7: Provide a brief bio for each investigator in the project *(less than one page each)*

Attachments
Please attach detailed budget and justification.
Appendix 2

Accelerating Medicines Partnership®
Bespoke Gene Therapy Consortium Principles

Publications
This project will operate under a "team science" approach, and publications generated by the Steering Committee will have joint authorship. Specific publication strategies will be developed by the Steering Committee prior to project start, including proposal for lead authors and co-authors. All publications and data resources generated by recipients of BGTC funding must acknowledge BGTC investigators and/or funder(s) and be made publicly accessible within 6 months of publication. Specific publication strategies will be discussed with the BGTC Steering Committee as needed.

Intellectual property (IP)
Given its precompetitive research focus and commitment to making results of that research available as broadly and promptly as possible, it is not expected that BGTC will generate novel IP. BGTC research partners may use pre-existing IP of the other BGTC research partners for work done under the partnership. BGTC research partners agree not to file patent applications on research discoveries made under the partnership, except in the rare instance when a consensus of FNIH and the BGTC agree that it is in the best interests of the partnership and public health to do so. IP developed under NIH awards are subject to applicable Federal law, regulation, and policies.