

**Bid Submission Form**  
**Participation in Assignment of Particle Concentration**  
**RFP 8.0**

Please complete the following fields:

*Contact Information – RFP 8.0*

*Contact Individual:	Mark A. Bowe, Ph.D.
Institution:	Genetic Therapy, Inc.
Address:	9 West Watkins Mill Road Gaithersburg, MD 20878
Phone Number:	301-258-4783
Fax Number:	301-330-2108
Email Address:	mark.bowe@pharma.novartis.com

**\*If laboratories are submitting a proposal as a group, a main contact should be provided along with contact information for each participating laboratory (attach additional copies of this form).**

Please indicate if your institution is also submitting proposals for the other activities:

- Determination of Infectious Titer
- Short-term/Field Stability Studies
- Long-term Stability Study
- Other Characterization
- Donation of Supplies/Other Services for Characterization Phase

**Bid Submission Form**  
**Participation in Assignment of Particle Concentration**  
**RFP 8.0**

General Capability Statement

Genetic Therapy, Inc. (GTI) proposes to participate in the assignment of particle concentration, using the OD<sub>260</sub> assay protocol provided by the ARMWG. GTI has been producing and analyzing adenoviral vectors for over ten years. The ARMWG's Standard Operating Procedure for Determination of Particle Concentration via Spectrophotometric Analysis will be performed by laboratory personnel of GTI's Core Technologies/Cell Biology Core. The Cell Biology Core has been analyzing adenoviral vectors under GLP conditions for three years (a qualified OD<sub>260</sub> assay for particle concentration is routinely used). The data from and reports of the analysis of the Adenoviral Reference Material will be reviewed minimally by one Cell Biology Core supervisor who did not participate in the assay, and by GTI Quality Assurance.

Personnel

GTI Cell Biology Core personnel are fully qualified to perform the ARMWG's Standard Operating Procedure. All laboratory personnel have a minimum of a bachelor's degree in Biology (or equivalent field). In addition, all laboratory personnel are trained in GLPs, biosafety, and specific laboratory techniques. A training file is maintained for each GLP employee.

Quality Assurance auditors have a minimum of a bachelor's degree in Biology (or equivalent field), a working knowledge of laboratory procedures, and are trained in auditing GLP and GMP documentation.

Key Personnel:

Cell Biology Core Manager: Mark A. Bowe, Ph.D. in Pharmacology (1992); employed by GTI since 9/1999.

Cell Biology Core/Lab Services Manager: Irina Burimski, M.S. in Cytology and Histology (1980); employed by GTI since 3/1990.

Quality Assurance Technical Specialist: Michael Stefanski, M.S. in Plant Virology (1988), M.S. in Biotechnology (1997); employed by GTI since 8/1997.

## Equipment

Cell Biology Core members have been trained on all equipment required by the protocol. Calibration of relevant equipment is as follows:

1. Beckman UV-Vis spectrophotometer, model DU 640: certified annually.
2. Gilson microliter pipettes: calibrated twice a year.
3. Pharmacia Biotech pH meter: calibrated daily before use with certified standards.
4. Scaltec Balance, model SBA 51: calibrated annually.

Equipment is maintained and used according to Standard Operating Procedures. Equipment logs and results of assays are reviewed by a Cell Biology Core supervisor.

## Timing

GTI is ready to accept Reference Material samples for testing as soon as they are available in mid to late September, 2001. The total time expected to perform the assay and review and report the results is 2-3 weeks after receipt of sample.