

Gene therapy process development support to optimize AAV yield & quality from gene to GMP production

Leverage our novel EpyQ™ AAV production system or traditional triple transfection for scalable solutions across all AAV serotypes

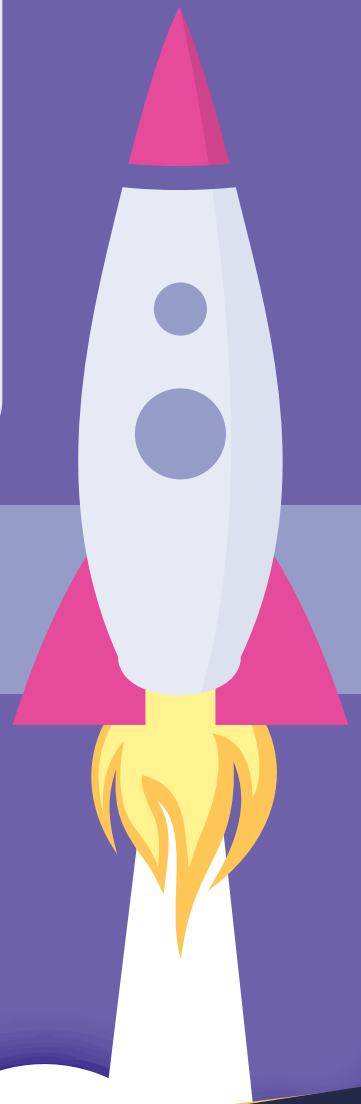
We help your team **Aim higher** with:

- A collaborative approach to delivering custom PD solutions quickly & efficiently
- Expertise in AAV development & manufacturing going back to 1991
- Global onboarding & tech transfer capabilities
- Risk analysis & mitigation for upstream & downstream optimization
- DoE-driven process optimization via extensive characterization with > 50 analytical assays
- Access to 15mL to 200+L bioreactors with high producing host cell options
- Innovative solutions for cell lysis & viral safety
- Feasibility & comparability studies from clinical to commercial
- CMC regulatory support & guidance from R&D to IND/IMPD submission

Mimicking our internal values, our process development services are driven by:

- 1 A quality-first approach to solutions tailored to your needs
- 2 Adaptability across clinical stages, scales & serotypes
- 3 Global network of labs aligned with FDA & EMA regulations

Aim
higher



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ADVANCED THERAPY MANUFACTURING SPECIALISTS