



Gene Therapy

Matthew Fuller, PhD

mfuller@ultragenyx.com

Vice President – Head of Gene Therapy Research

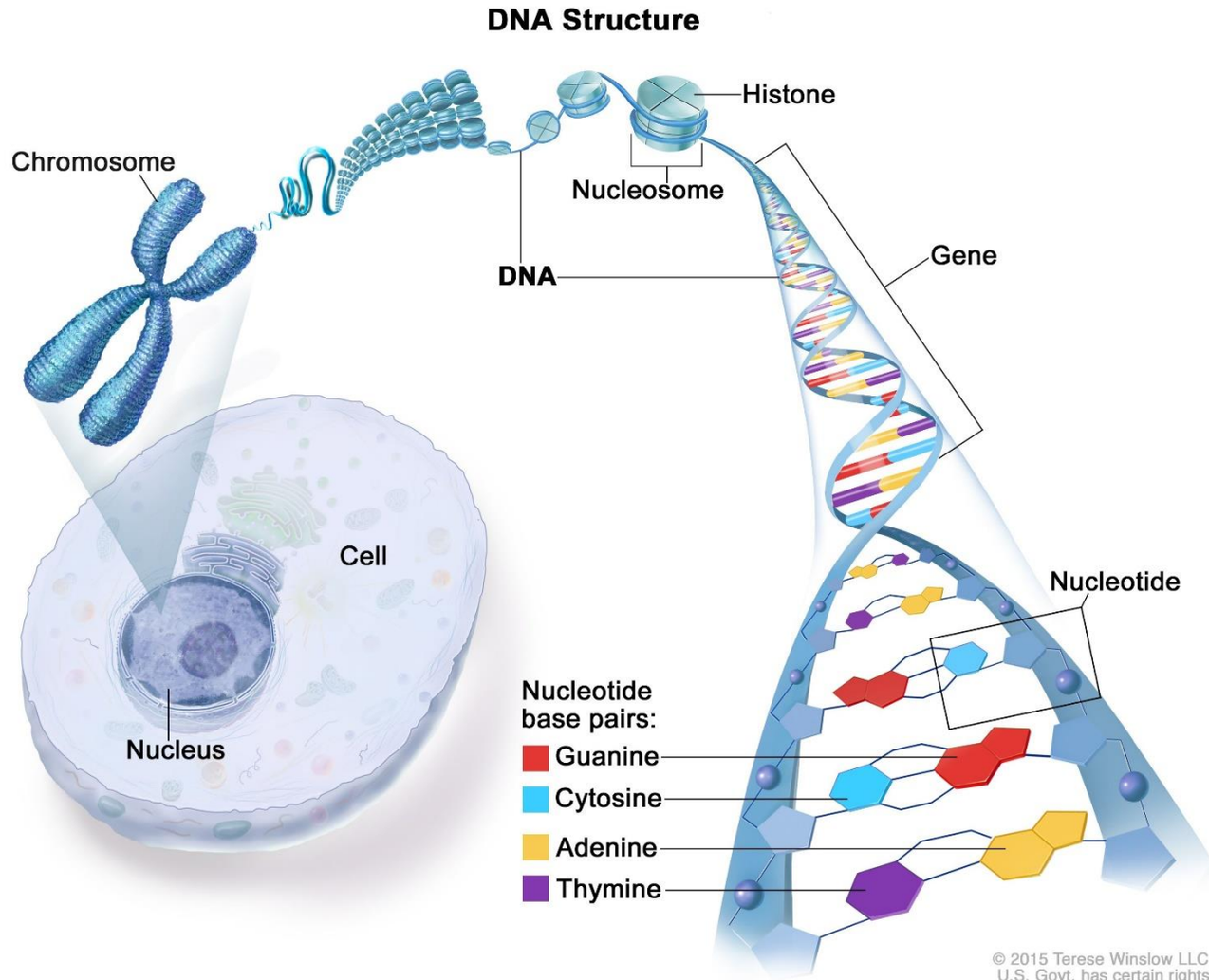
November 13th, 2024



Agenda

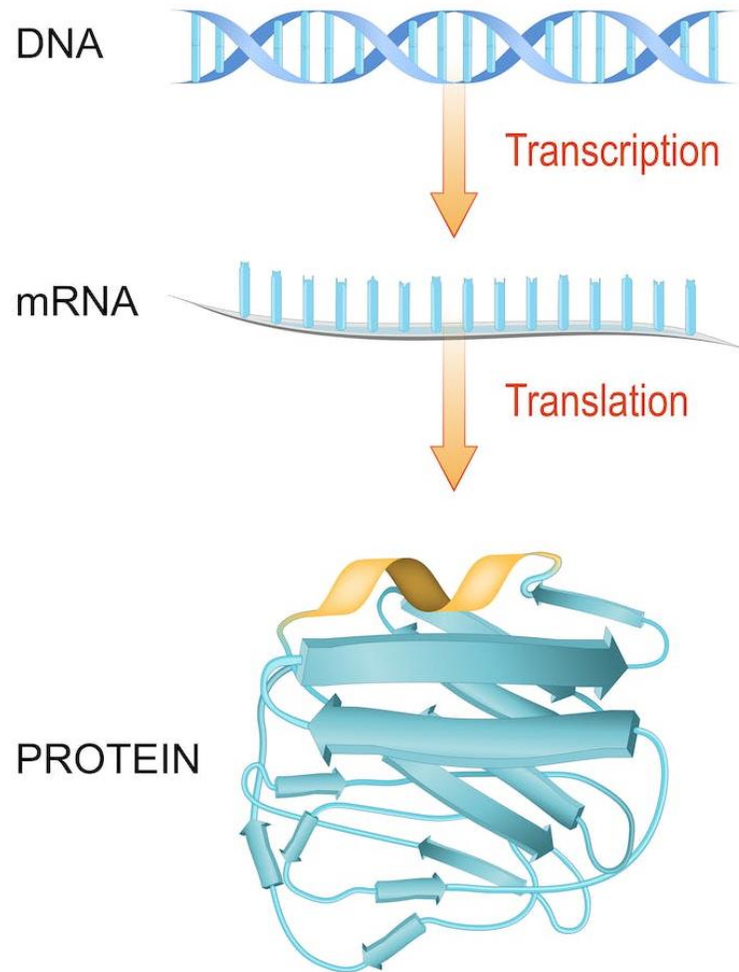
1. Background – Foundational Biology
2. Gene Therapy Approaches
3. Key Takeaways

Basic Architecture of Genes and DNA



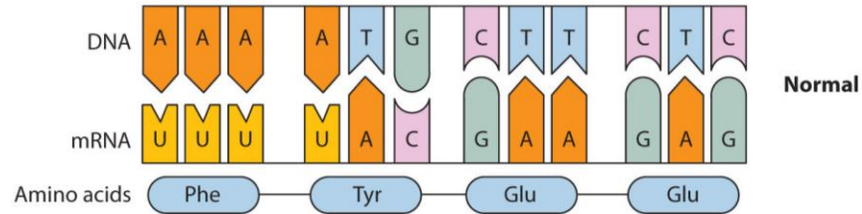
- **Gene** – A basic genetic unit encoding a function
 - Located in a specific section of DNA on a specific chromosome
 - Encoded by the DNA nucleotide “alphabet” – A, G, T & C
 - Genes encoding overt functions typically code for proteins
 - *Alterations* in genes can, but do not always, cause dysfunction
- **NPHP1 gene** is located on **Chromosome 2 [2q13]**
- **DARS2 gene** is located on **Chromosome 1 [1q25.1]**
- **KIZ gene** is location on **Chromosome 20 [20p11.23]**

DNA Encodes for Proteins with Specific Functions



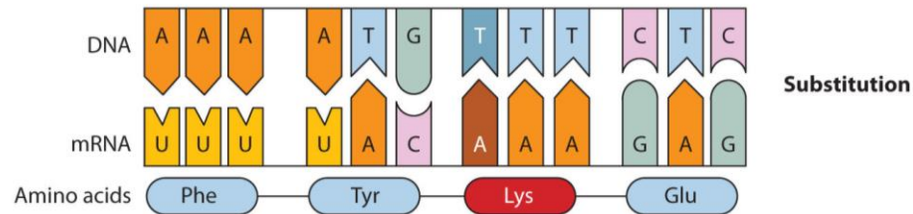
- DNA encodes ~20,000 genes with discrete cellular functions utilized by our cells & bodies for our day-to-day existence
- These functions are facilitated by pathways that utilize the 'DNA -> mRNA -> Protein' program to provide the necessary elements to carry out these day-to-day tasks
 - Examples of protein functions include processing of metabolites, muscle contraction, DNA replication, movement of other proteins, activation or inhibition of other proteins, and much more...
- Gene therapy seeks to replace or repair these specific pathways & molecules when typical function is not being provided due to mutation(s)

Common Types of DNA Mutations



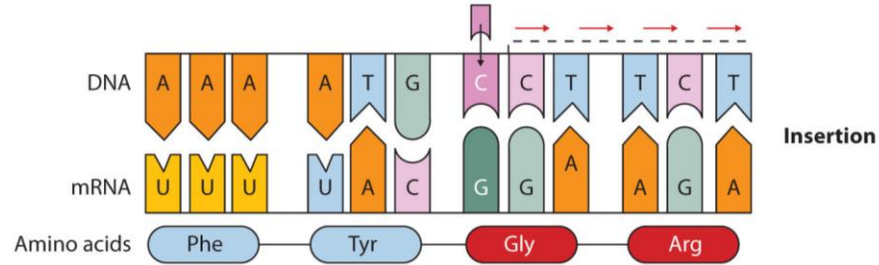
THE CAT ATE THE RAT
THE **K**AT ATE THE RAT

Silent Mutation
The meaning of the sentence is the same.



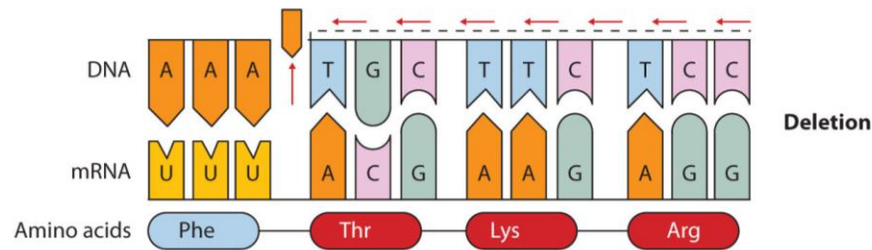
THE CAT ATE THE RAT
THE **H**AT ATE THE RAT

Substitution Mutation
The meaning of the sentence is changed.



THE CAT ATE THE RAT
THE **E**CA TAT ETH ERA T

Insertion Mutation
The sentence no longer makes sense.

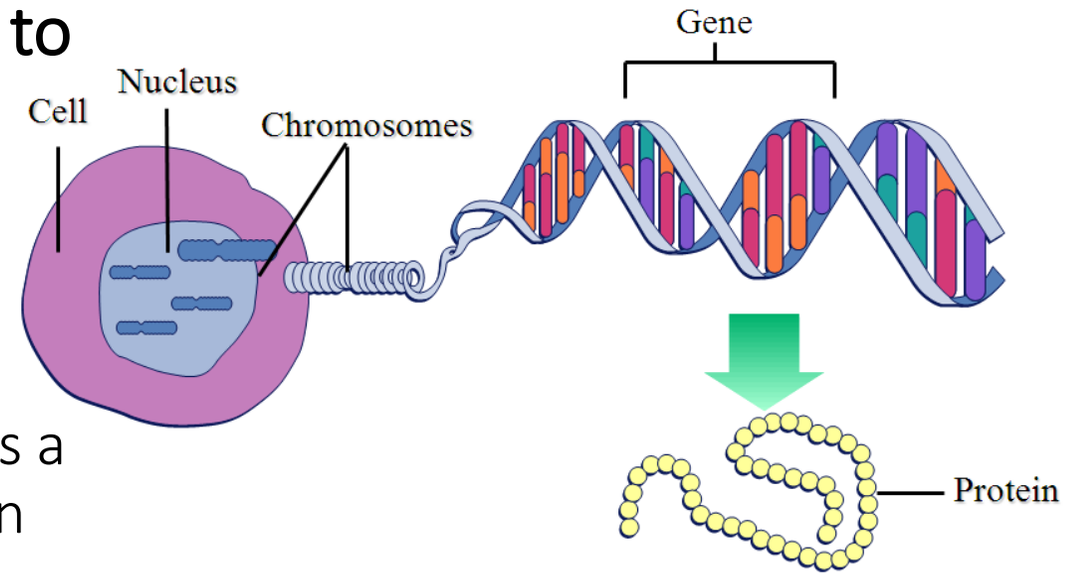


THE CAT ATE THE RAT
THE **C**A TET HER AT

Deletion Mutation
The sentence no longer makes sense.

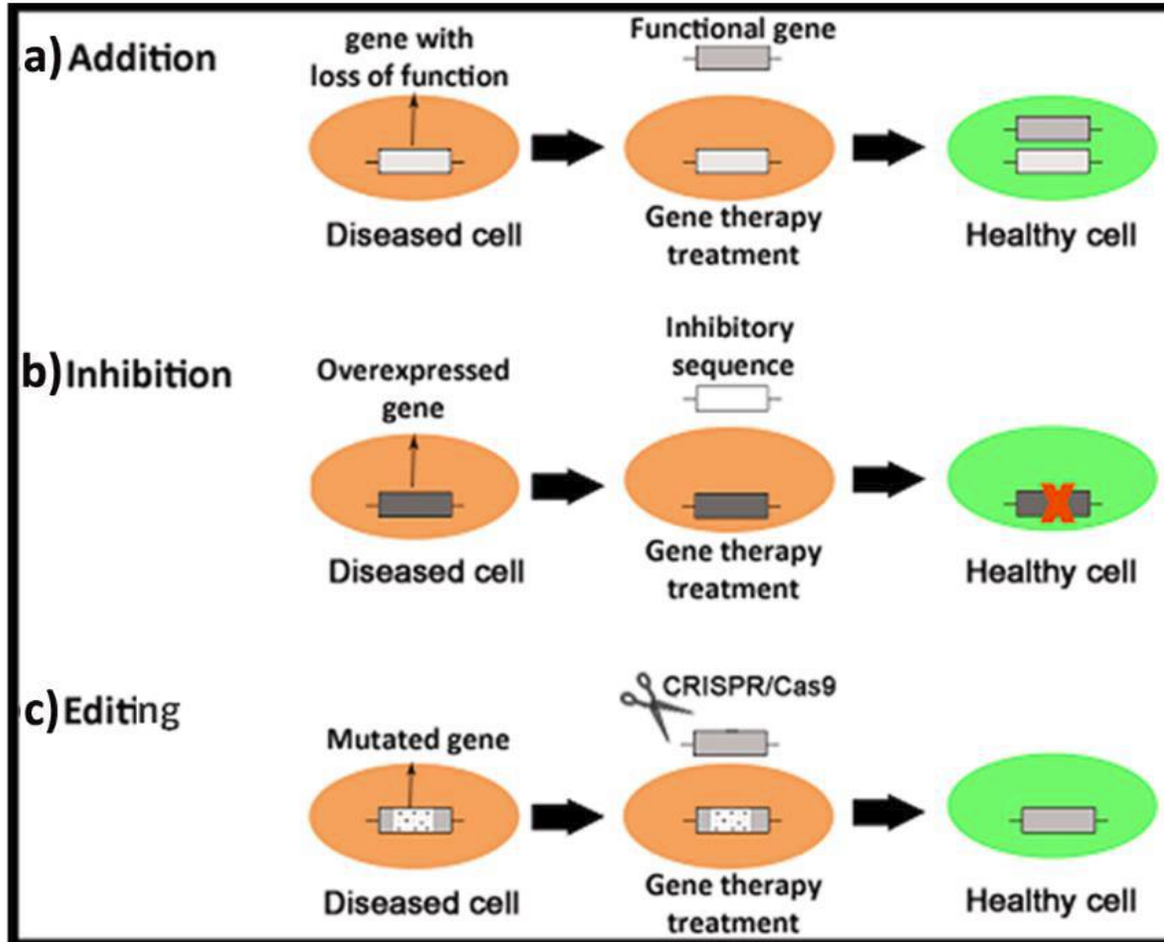
Basic Definition of Gene Therapy

- Treatment or prevention of a [genetic] disease *via* introduction of genetic material expected to provide a necessary function
- First developed in 1972 when Theodore Friedmann and Richard Roblin published a paper in *Science* called "Gene therapy for human genetic disease?"
- The first patient to be treated with gene therapy was a four-year-old girl treated at the NIH Clinical Center in 1990
 - She had a congenital disease called adenosine deaminase (ADA) deficiency which severely affects immunity and the ability to fight infections
 - Treatment was considered successful (*ex vivo*)



<http://igbiology.blogspot.com/2014/03/chromosomes-dna-genes-and-alleles.html>

Gene Therapy Approaches



- Gene therapy approaches include the delivery of a package/cargo designed to:
 - Add back a functional or wildtype copy of a mutated or missing gene that is causing disease
 - Inhibit, inactivate, or “knock out,” a mutated or overexpressed gene that is functioning improperly
 - Edit a mutated gene back to a functional or wildtype copy of that gene

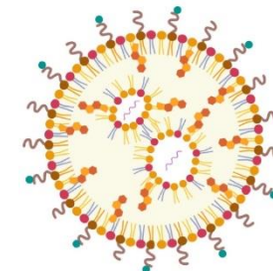
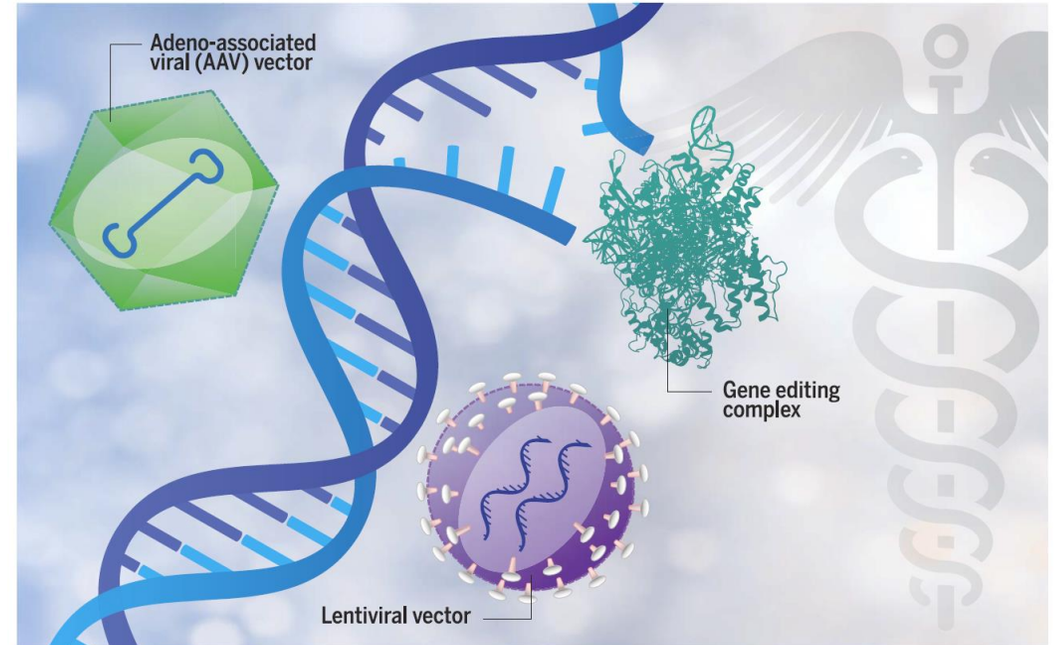
Int. J. Mol. Sci. 2021, 22, 7545. <https://doi.org/10.3390/ijms22147545>

Note – “wildtype” refers to the natural or common version of a specific protein, DNA, virus, etc. that exists in nature

Common Tools of Gene Therapy

■ The 3 most basic tools currently utilized for Gene Therapy are:

- Adeno-associated virus (AAV)
 - Delivery Vehicle and Cargo
- Lentivirus
 - Delivery Vehicle and Cargo
- Gene Editing complex
 - Cargo
 - Must be coupled with Delivery Vehicle
- *mRNA*
 - Delivery via Lipid Nanoparticles (LNPs)
 - *Genetic therapy*



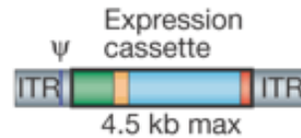
Lipid nanoparticle (LNP)

Extended Toolbox of Gene Therapy Vectors

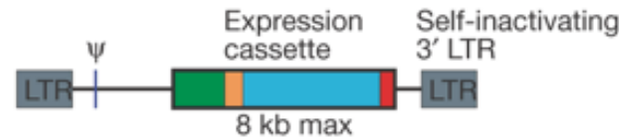
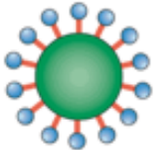
Adenovirus (~36 kb genome)



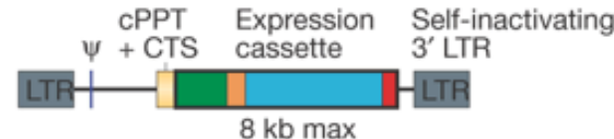
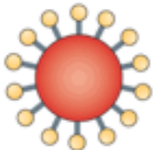
Adeno-associated virus (4.7 kb genome)



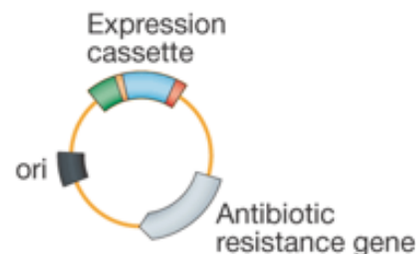
Retrovirus (7-10 kb genome)



Lentivirus (9-10 kb genome)



Liposome + plasmid (unlimited sized genome)



• Adenovirus

- Antiviral vaccines; Anticancer therapy; Larger transgene size; Potential immune response to vector

• AAV

- Ideal for targeting non-dividing cells; Ideal for *in vivo* delivery; Ideal for gene replacement; Smaller transgene size

• Retrovirus

- Require active cell division for infection (target dividing cells); Lower safety profile than lentivirus due to higher rate of oncogenesis

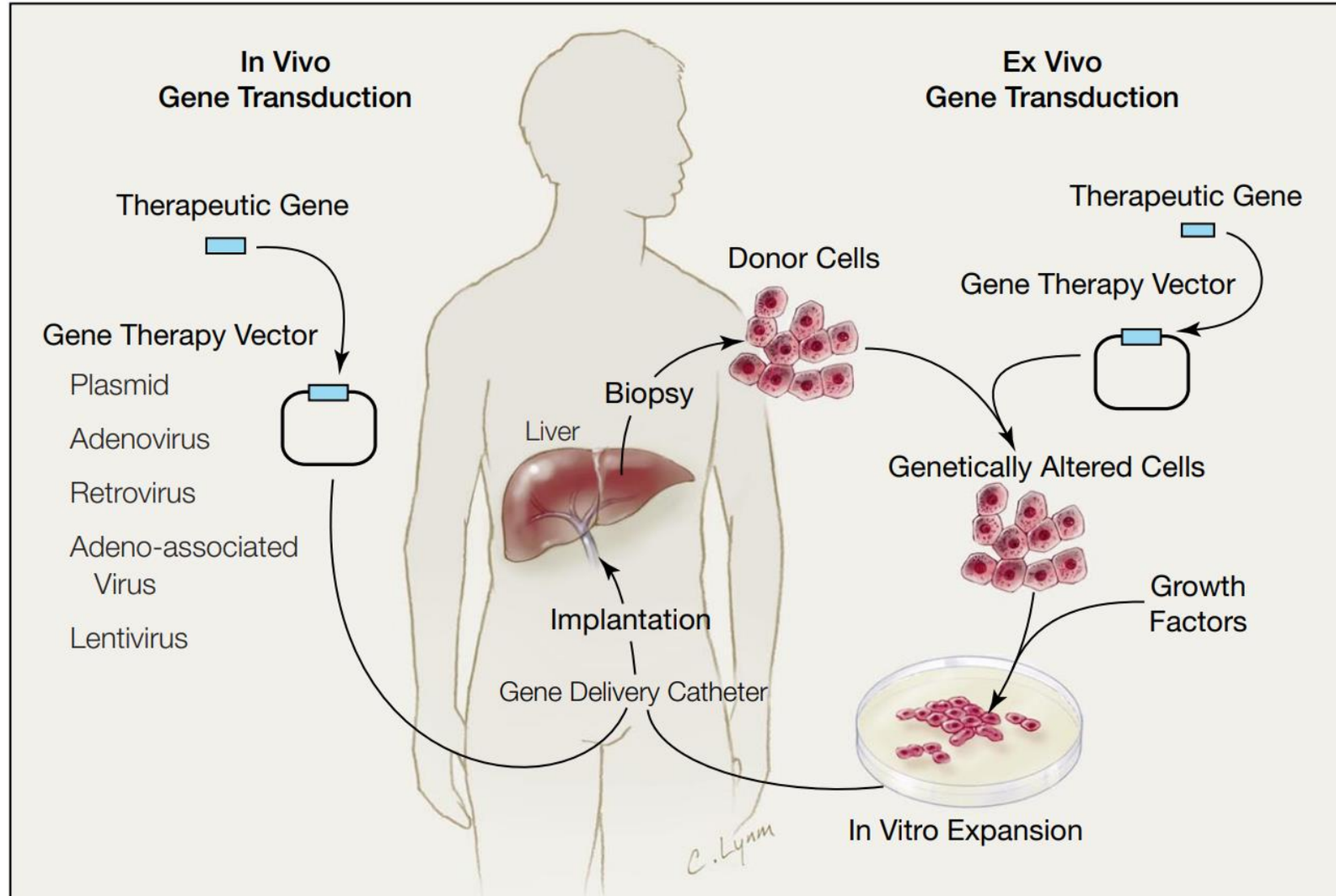
• Lentivirus

- Ideal for targeting non-dividing cells; Ideal for *ex vivo* delivery; Larger transgene size than rAAV, less than Adenovirus

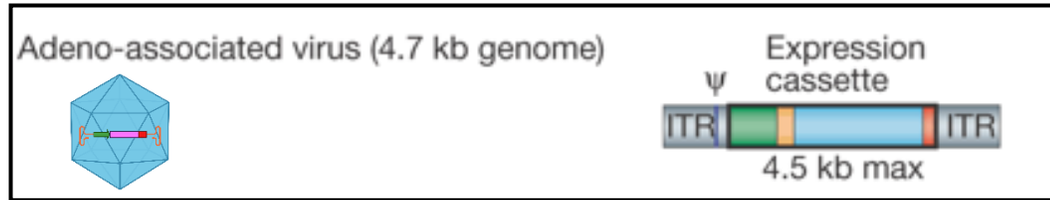
• Liposome + Plasmid

- Very large transgene size; Complicated manufacturing and delivery

In Vivo versus Ex Vivo Gene Delivery Requirements Influence Vector Choice



Primary Viral Tools of Gene Therapy – AAV & Lentivirus

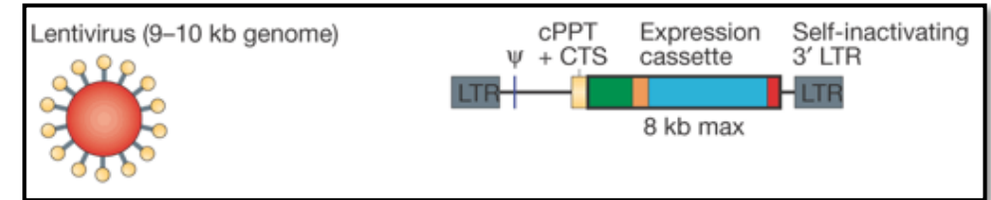


• Advantages

- Ideal for *in vivo* delivery (directly into patient)
- Ideal for gene replacement
- Can be used for CRISPR, but small genome size limitations exist
- Very low integration rates
- Multiple capsid choices tailor cellular tropism
- Academic and industry manufacturing experience

• Challenges

- Small genome size (≤ 5 kilobases)
- *Expensive* to manufacture
- Single dose limitation*
- Immune reactions to treatment in small number of patients not well understood



• Advantages

- Ideal for *ex vivo* delivery (into isolated patient cells, which are then returned to patient)
- Ideal for targeting nondividing cells
- Infect wide variety of cell types
- Larger transgene size than rAAV (Lenti ≈ 9 kilobases), smaller than Adenovirus (~ 36 kilobases)

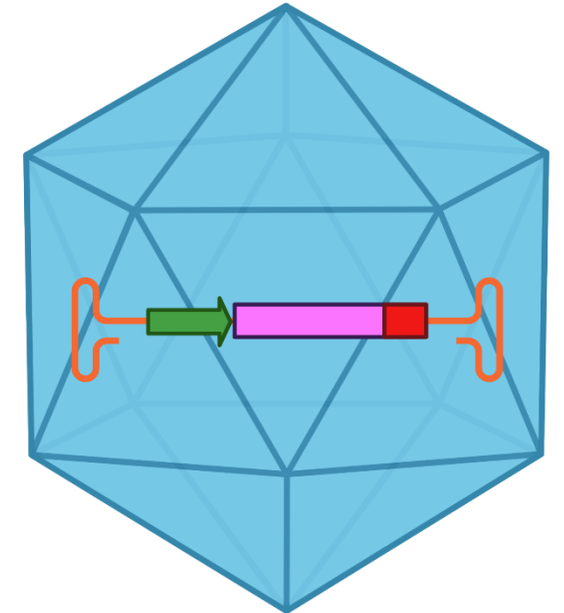
• Challenges

- Lentiviral integration poses higher oncogenic/genotoxic risk than AAV
- Potential activation of neighboring genes post-integration

* The field is attempting to address this with IdeS
(<https://www.nature.com/articles/s41591-020-0911-7>)

What Makes a “Good Candidate” for *Addition* Gene Therapy?

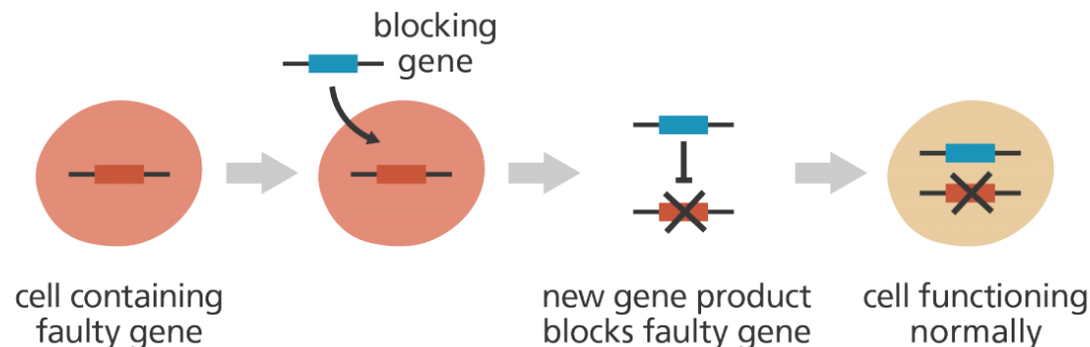
- Does your transgene fit?
 - AAV maximum packaging capacity is ~5 kilobases of DNA
 - Lentivirus maximum packaging capacity is ~9 kilobases DNA
 - Note – This maximum capacity is not for transgene alone, but requires consideration of *required* viral and regulatory elements as well (used to turn gene expression on, package into virus, terminate mRNA)
- Can the delivery vehicle (AAV/Lentivirus) infect and transfer genetic material to the target cell(s) important for correcting the disease?
- Are the target cell(s) amenable to the gene therapy approach/cargo you are using? (addition vs. inhibition)
 - Important to consider gene expression levels associated with a disease phenotype, especially in CNS-related diseases [Goldilocks scenario]



AAV Capsid

What Makes a “Good Candidate” for *Inhibition* Gene Therapy?

- Is there a well understood target to inhibit?
 - This target can be direct (e.g., the protein causing the disease) or...
 - This target can be indirect (e.g., a genetic component modifying the protein causing the disease)
 - Examples include expression of an inhibitory protein, an inhibitory sequence or a ‘suicide gene’ to kill affected cells (cancer or infectious disease)
 - Long term delivery methods (e.g., AAV) mostly used for infectious disease and cancer
- Can the delivery vehicle reach and transfer inhibitory genetic material to the target cell(s) important for correcting the disease?
- Are the target cell(s) amenable to the gene therapy approach/cargo you are using?



Why is Gene Therapy a Compelling Therapeutic Modality?

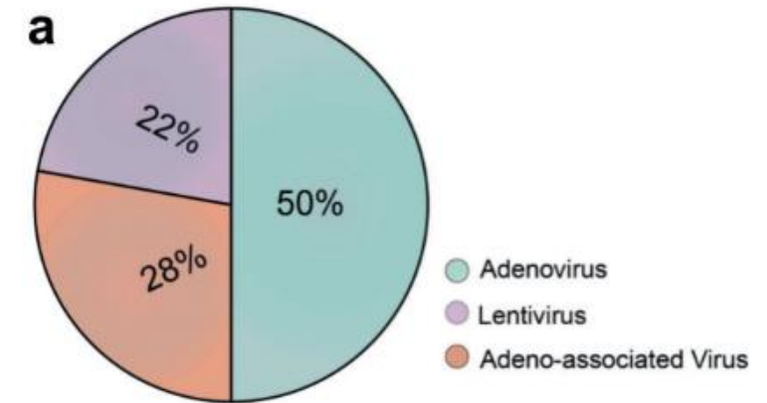
- *Potential* for halting, treating *or* curing* disease using a one-time** treatment
- Broad experience in global clinical trials
- Increased awareness and acceptance due to recent product approvals and compelling late-stage clinical results
- Regulators and payers becoming more familiar with therapeutic paradigm

* The definition of 'cure' is a complicated, moving target

** Limited by lifetime/dilution of treated cells

Success Stories of Gene Therapy

- >1500 GT trials have been initiated across the 3 most common delivery vehicles
- 24 approved Gene Therapy products; 7 approved Adeno-associated Virus (AAV) products; 7 approved Lentivirus products; 1 approved CRISPR product (for 2 indications);
 - Luxterna (Spark/Roche) – Leber’s congenital amaurosis (RPE65 deficiency)
 - Cost = \$850,000, at launch
 - Zolgensma (AveXis/Novartis) – Spinal Muscular Atrophy (SMN1)
 - Cost = \$2.125M, at launch
 - Hemgenix (CSL Behring/UniQure) – Hemophilia B (FIX)
 - Cost = \$3.5M, at launch
 - Roctavian (BioMarin) = Hemophilia A (FVIII)
 - Cost = \$2.9M, at launch
 - Upstaza (PTC Therapeutics) – Aromatic l-amino acid decarboxylase deficiency
 - Cost = \$3.7M, at launch
 - Elevidys (Sarepta) – Duchenne Muscular Dystrophy (DMD) [≥4 years-old]
 - Cost = \$3.2M, at launch
 - Beqvez (Pfizer) – Hemophilia B (FIX)
 - Cost = \$3.5M, at launch
 - Casgevy (CRISPR Therapeutics/Vertex) – Sickle Cell Disease and transfusion-dependent Beta-thalassemia [≥12 years old]
 - Cost = \$2.2M, at launch



b

| Vectors | Number of clinical trials |
|------------------------|---------------------------|
| Adenovirus | 575 |
| Adeno-associated Virus | 250 |
| Lentivirus | 315 |
| Total | 1140 |

Signal Transduction and Targeted Therapy (2021)6:53

Keys to Recent Successful Gene Therapy Outcomes

- Solid basic and clinical science
- High unmet medical need
- Good cellular/tissue target choice
- Increased platform understanding by Health Authorities
- **What limitations exist?**
 - Commercial CMC
 - Continued stringency of Health Authority CMC requirements
 - Extension of clinical success to new targets
 - Safety concerns with high dose AAV delivery ($>1-2 \times 10^{14}$ vg per kg)
 - SMA – AAV9
 - XLMTM – AAV8
 - DMD – AAV9
 - Continued study of immunological reactions to gene therapy

Key Takeaways

- Gene therapy has the *potential* for halting, treating *or* curing* disease using a one-time** treatment
 - There are multiple successful, FDA approved examples *via* recombinant AAV and Lentiviral vectors
- Success is *critically dependent* upon solid basic and clinical science knowledge, a targetable cellular/tissue target choice, and the ability to manufacture the vector
- Each gene therapy strategy and delivery vehicle has pros and cons
 - Strategy and delivery vehicle must be matched to the biology of the disease and target cells
- Gene therapy manufacturing and reimbursement are *expensive*
 - The field is continually working to improve manufacturability and reduce costs
 - Broader adoption of gene therapy treatments and continued understanding of treatment paradigm by regulators and payers should lead to reduced costs over time



Sponsored by Ultragenyx

Thank You





Sponsored by Ultragenyx





Sponsored by Ultragenyx

Appendix

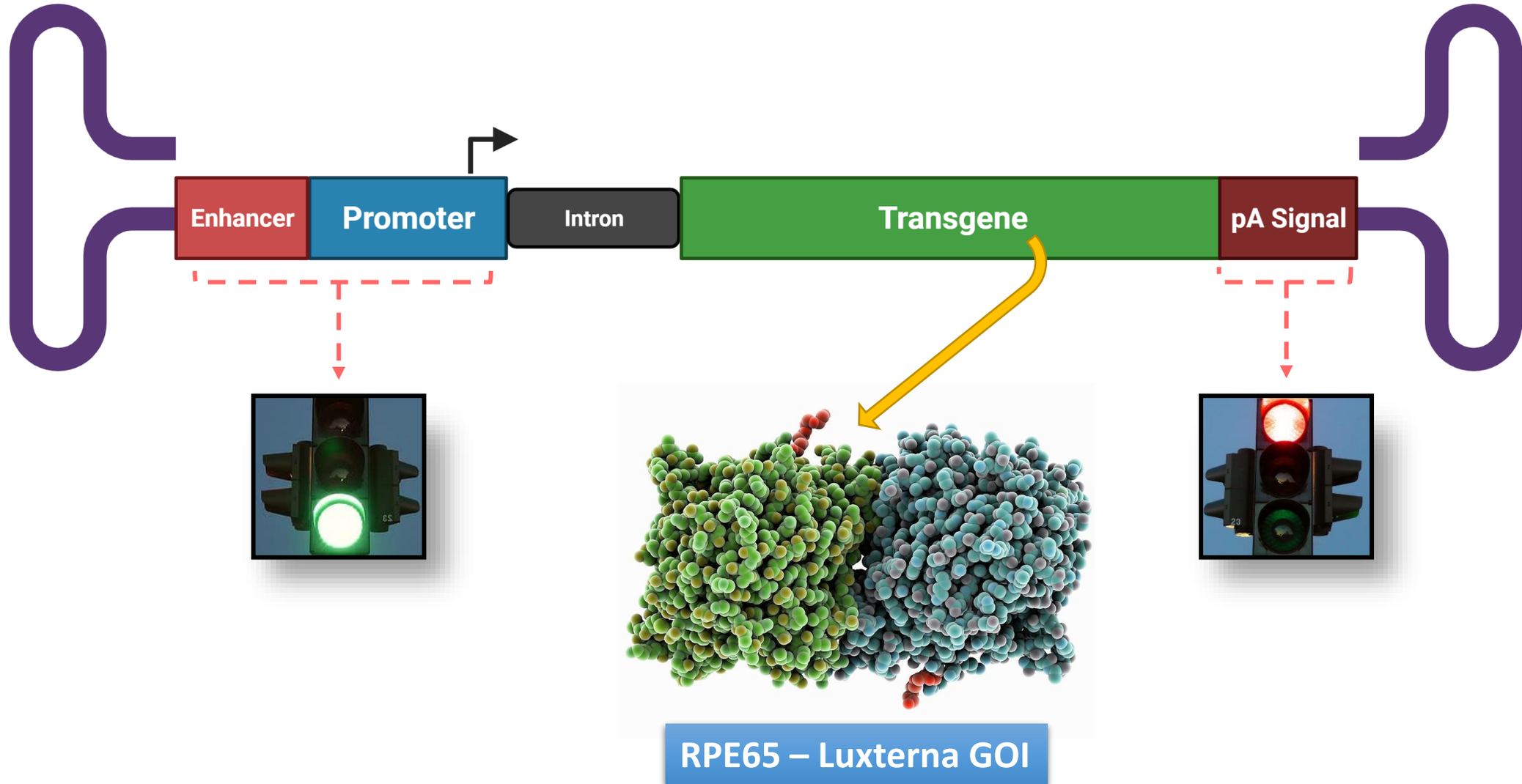




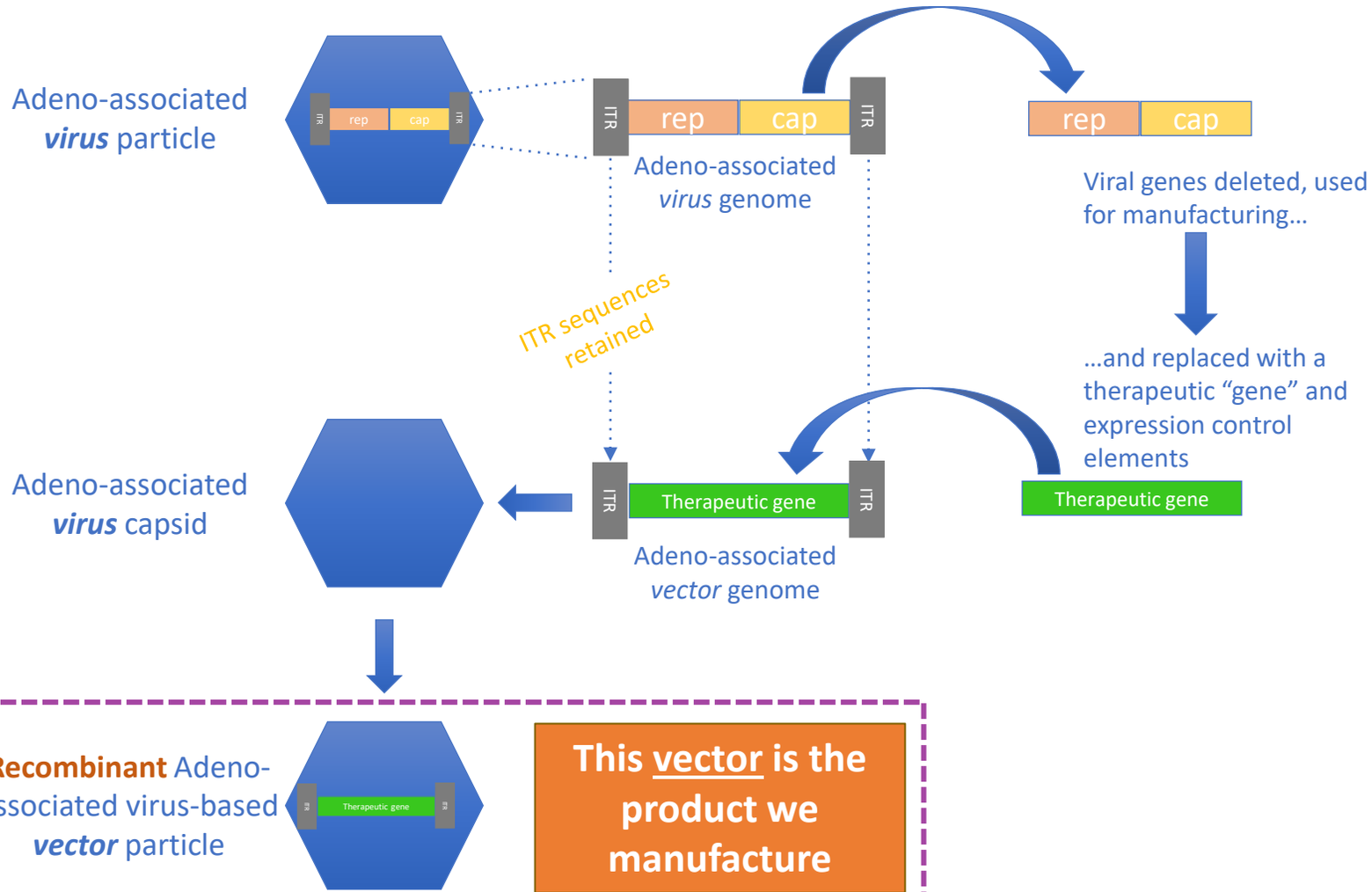
RARE ENTREPRENEUR
BOOTCAMP

How to Design and Generate an AAV GT Vector

Fundamental Design of a Recombinant AAV



Basic recombinant AAV Molecular Engineering



Step 1: "Gut" wildtype AAV virus

Step 2: Insert gene of interest (GOI)

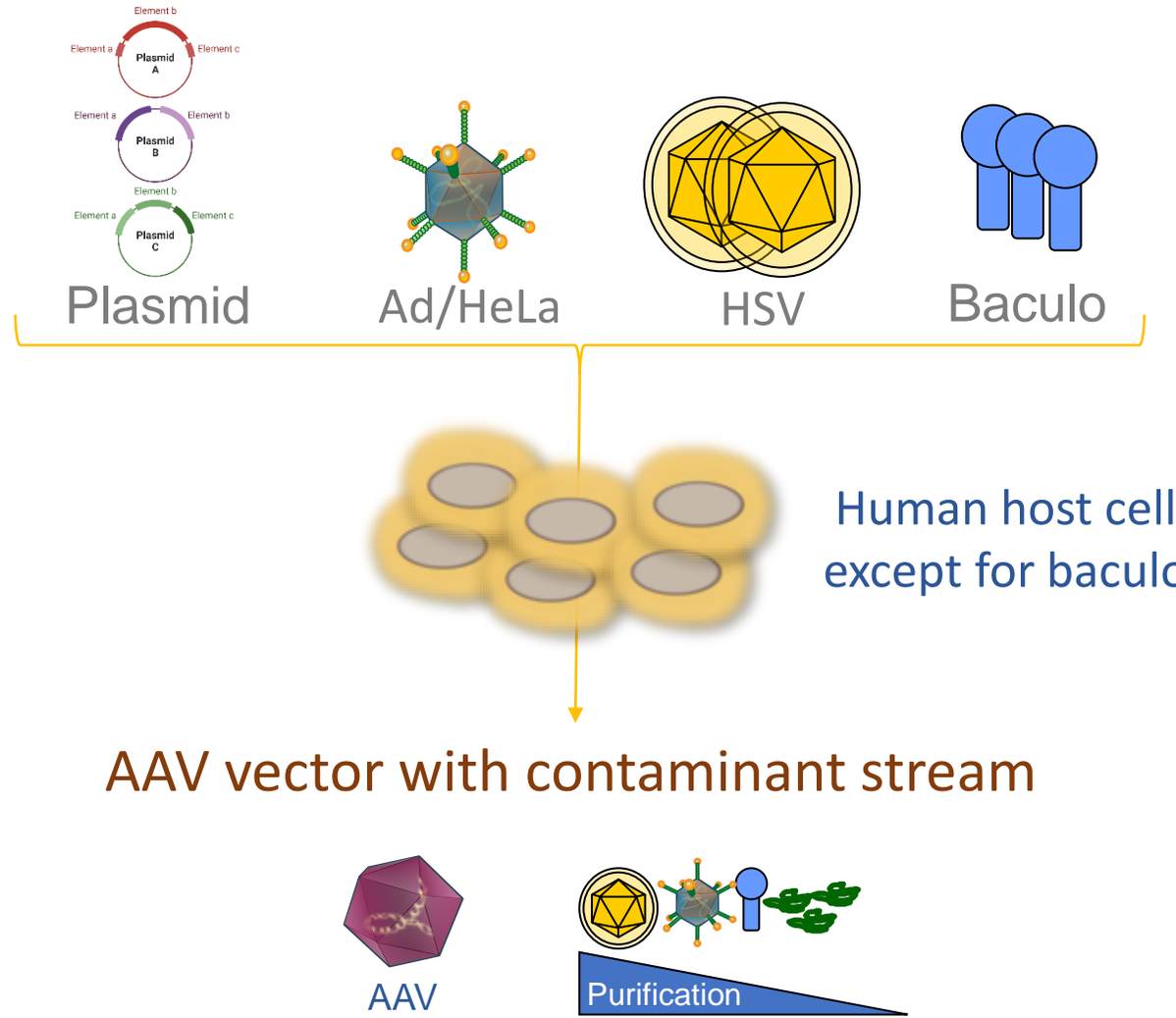
Step 3: Manufacture recombinant AAV

AAV Vector Manufacturing – Overview

AAV rep+cap functions and helper functions must be provided

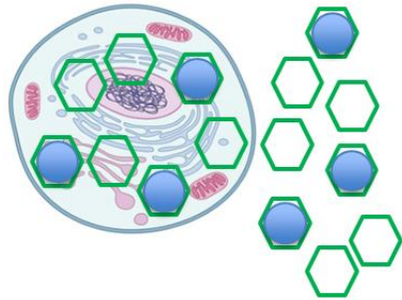
Commercial success requires bioreactor production platform and scale

Scalable operations must be designed to purify the product at acceptable yield and quality, while removing contaminants

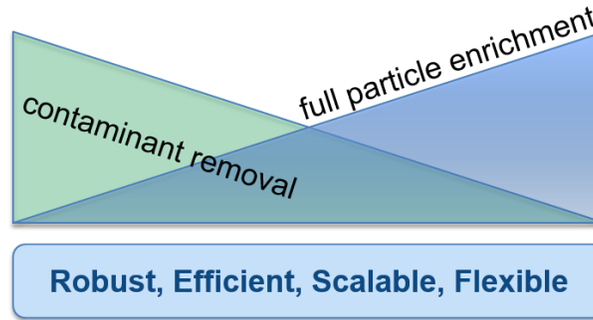


AAV Vector Purification

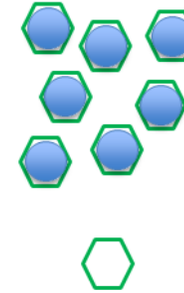
Upstream Cell-based
Platform



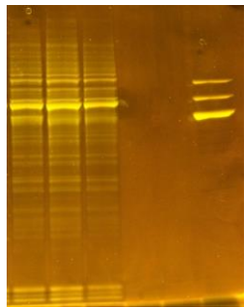
Downstream Purification
Platform



Final Product
*highly enriched for active
DNA containing vector*

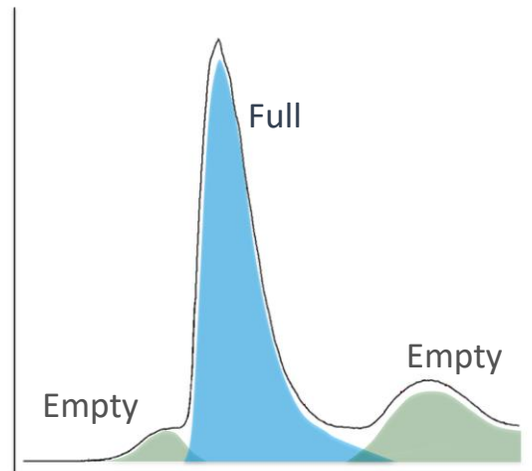


Affinity Capture

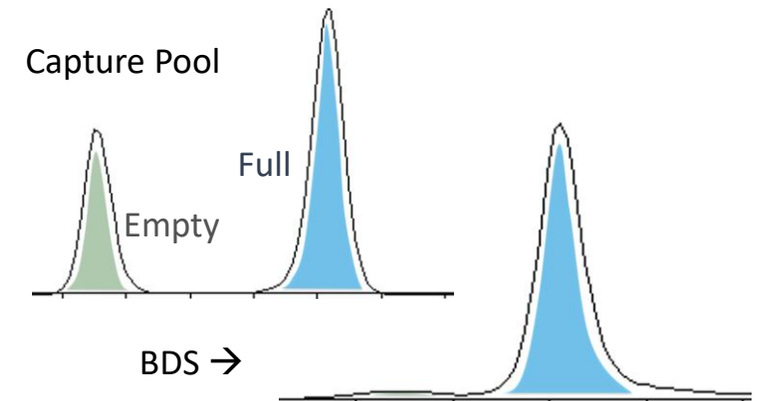


Load/Wash | Pool

Polishing Step



Analytical Ultracentrifuge



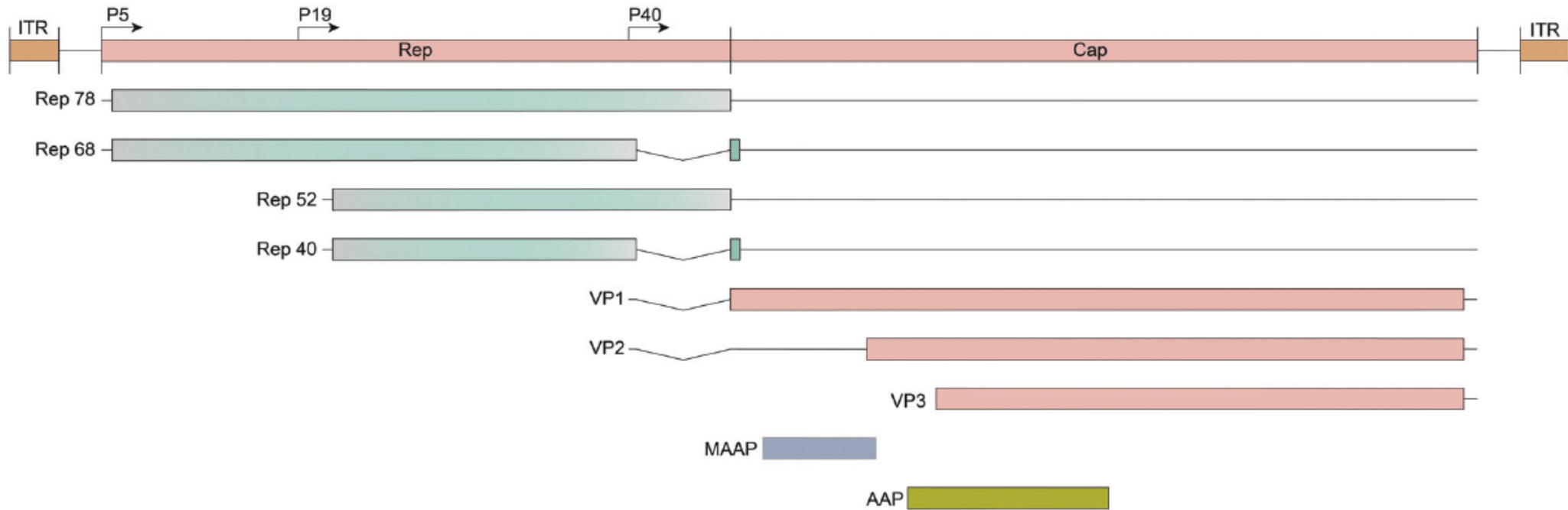
- **Gene therapy has the *potential* for halting, treating or curing disease using a one-time* treatment**
 - There are successful, FDA approved examples *via* recombinant AAV and Lentiviral vectors
- **Success is *critically dependent* upon solid basic and clinical science knowledge, a targetable cellular/tissue target choice, and the ability to manufacture the vector**
- **Each gene therapy strategy and delivery vehicle has pros and cons**
 - Strategy and delivery vehicle must be matched to the biology of the disease and target cells
- **Gene therapy manufacturing and reimbursement are *expensive***
 - The field is continually working improve manufacturability and reduce costs
 - Broader adoption of gene therapy treatments and continued understanding of treatment paradigm by regulators and payers should lead to reduced costs over time



RARE ENTREPRENEUR
BOOTCAMP

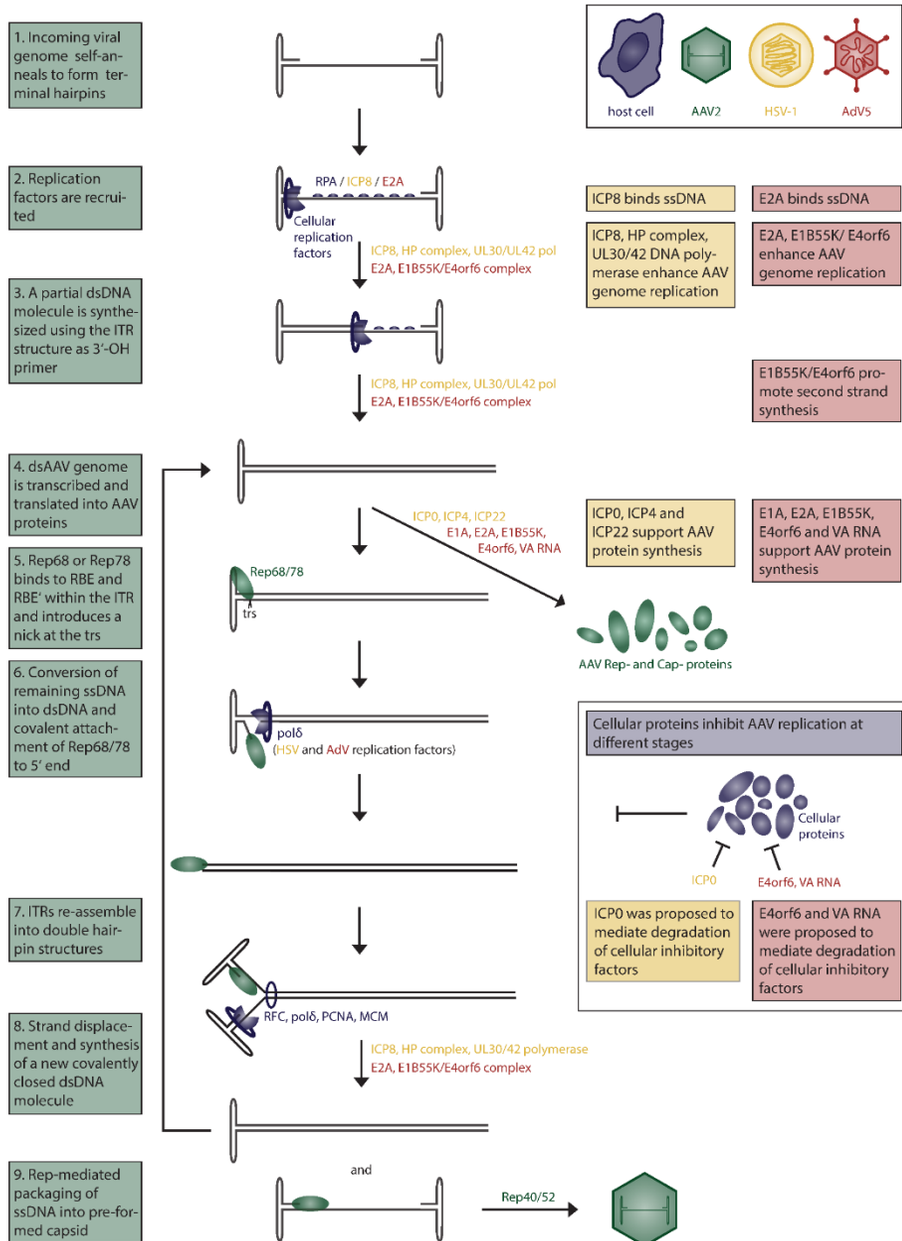
A bit more detailed information on
AAV

AAV Protein Expression



- AAV “Large Rep” proteins provide a nicking function required for genome replication [Rep78 and Rep68]
- AAV “Small Rep” proteins are involved in packaging the genome into the preformed capsid [Rep52 and Rep 40]
- AAV Capsid proteins (“Cap”) form the structure that the genome is packaged into and is the delivery mechanism of gene therapy [VP1, VP2, VP3]

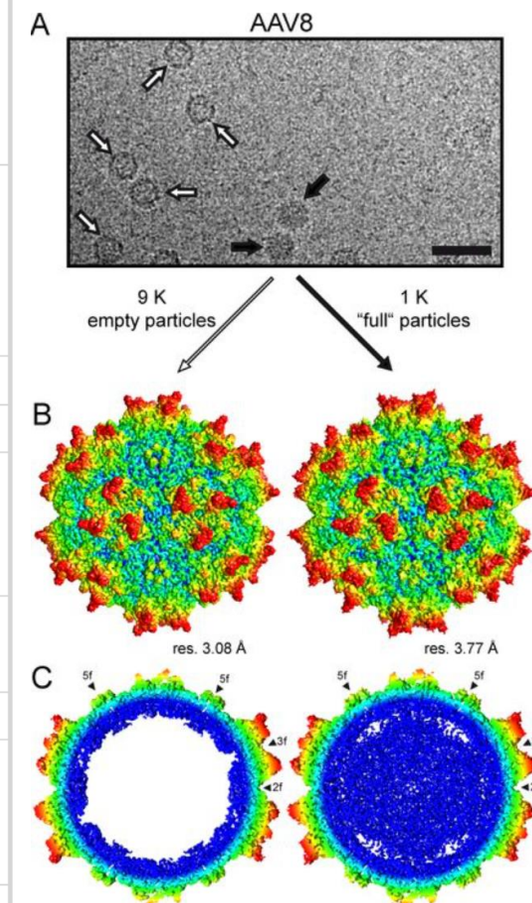
AAV Replication – Helper Virus



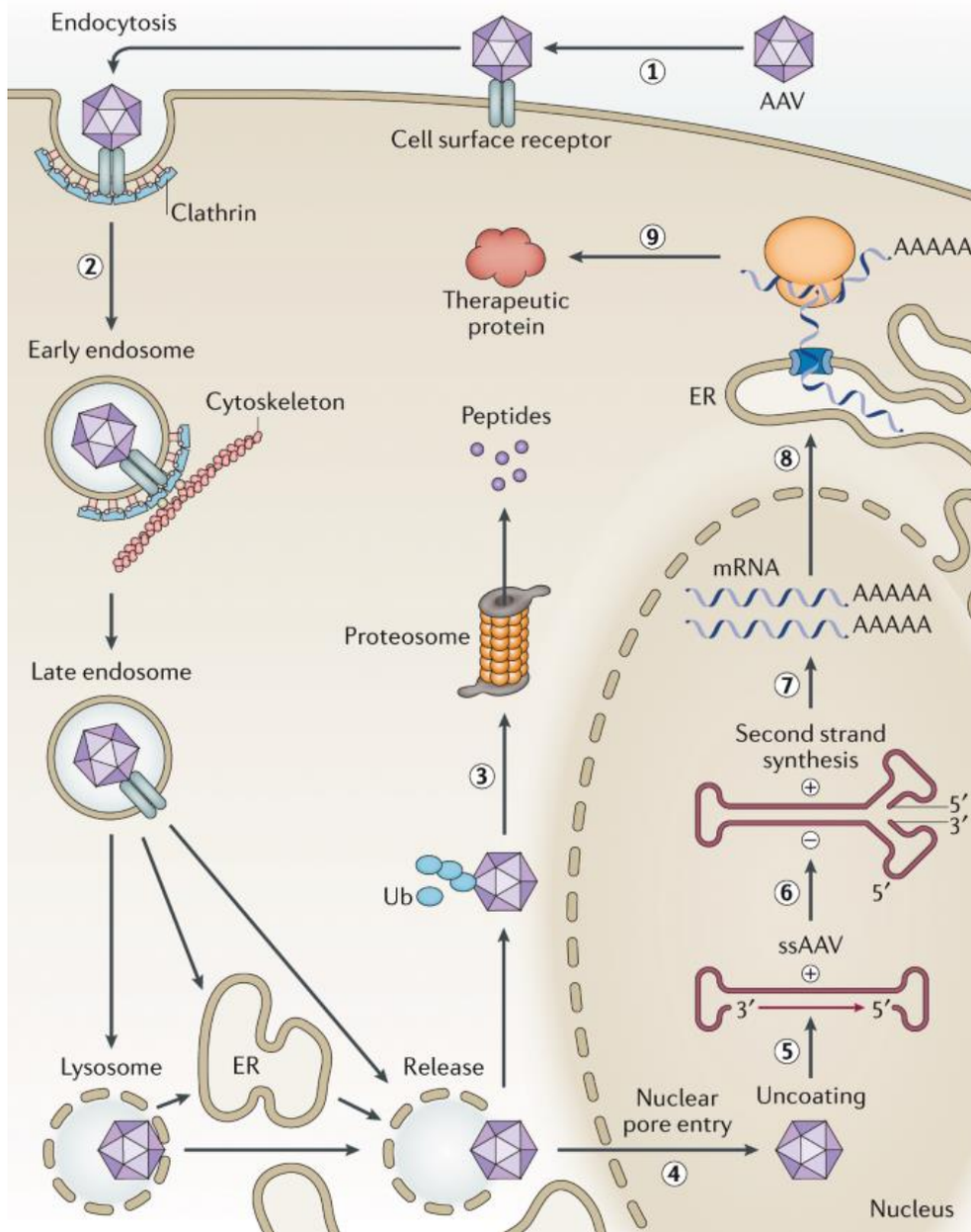
- AAV co-opts both cellular and viral factors to facilitate completion of its life cycle (genome replication and packaging)
- Cellular factors are used to replicate the AAV genome
- Helper virus factors are used to support replication of the AAV genome
- AAV Rep proteins are used to facilitate replication and packaging of the AAV genome
- AAV Cap proteins associate to form the capsid

AAV Capsid Selection

| AAV serotype | Origin of isolation | Primary receptor | Co-receptor | Tissue tropism | Condition (ClinicalTrials.gov identifier) | Approved drug |
|--------------|---------------------|----------------------|----------------------------------|------------------------------|---|---|
| AAV1 | Monkey | Sialic acid | AAVR | Muscle, CNS, heart | Muscle diseases (NCT01519349) | None |
| | | | | | Heart failure (NCT01643330) | |
| | | | | | AAT deficiency (NCT01054339, NCT00430768) | |
| AAV2 | Human | Heparin | Integrin, FGFR, HGFR, LamR, AAVR | Liver, CNS, muscle | Eye diseases (NCT00643747) | Luxturna for Leber congenital amaurosis |
| | | | | | Haemophilia (NCT00515710) | |
| | | | | | CNS diseases (NCT00400634) | |
| | | | | | AAT deficiency (NCT00377416) | |
| AAV3 | Human | Heparin | FGFR, HGFR LamR, AAVR | Muscle, stem cells | No trials underway | None |
| AAV4 | Monkey | Sialic acid | Unknown | Eye, CNS | Eye diseases (NCT01496040) | None |
| AAV5 | Human | Sialic acid | PDGFR, AAVR | CNS, lung, eye | Haemophilia (NCT03520712) | None |
| | | | | | Eye diseases (NCT02781480) | |
| | | | | | AIP (NCT02082860) | |
| AAV6 | Human | Heparin, sialic acid | EGFR, AAVR | Muscle, CNS, heart, lung | Haemophilia (NCT03061201) | None |
| | | | | | CNS diseases (NCT02702115) | |
| AAV7 | Monkey | Unknown | Unknown | Muscle, CNS | No trials underway | None |
| AAV8 | Monkey | Unknown | LamR, AAVR | Liver, muscle, pancreas, CNS | Eye diseases (NCT03066258) | None |
| | | | | | Haemophilia (NCT00979238) | |
| | | | | | Muscle diseases (NCT03199469) | |
| AAV9 | Human | Galactose | LamR, AAVR | Every tissue | CNS diseases (NCT02122952) | Zolgensma for spinal muscular atrophy |
| | | | | | Muscle diseases (NCT03362502) | |
| AAV10 | Monkey | Unknown | Unknown | Muscle | No trials underway | None |
| AAV11 | Monkey | Unknown | Unknown | Unknown | No trials underway | None |
| AAV12 | Human | Unknown | Unknown | Nasal | No trials underway | None |



AAV Transduction



- AAV capsid interacts with the external cellular receptor and is endocytosed
- AAV capsid interacts with internal endosomal receptor
- Endosome matures from early to late stage, pH change induces conformational change in AAV capsid externalizing VP1
- VP1 phospholipase (PLA) activity opens the endosome allowing capsid escape
- Capsid traffics to the nucleus, disassembles and releases the single-stranded DNA
- WT virus expresses viral proteins and replicates
- Recombinant AAV concatamerizes and resides as an episome loosely associated with cellular chromatin (forms circular DNA molecule)



RARE ENTREPRENEUR
BOOTCAMP

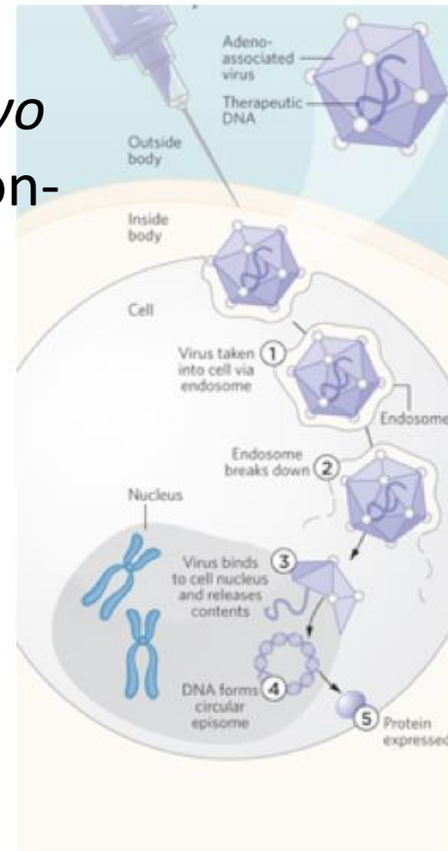
Transition Slide

Appendix

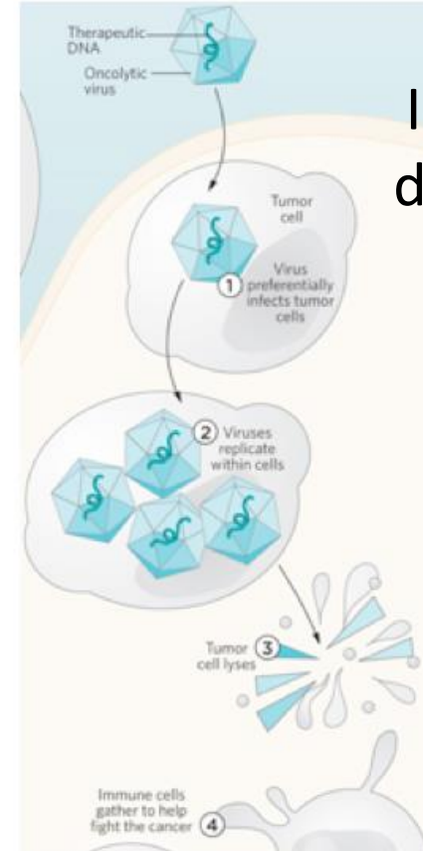
GT Overview: Introducing Genetic Material via Viruses

Ideal for *in vivo* delivery to non-dividing cell targets

Non-Integrating (AAV)



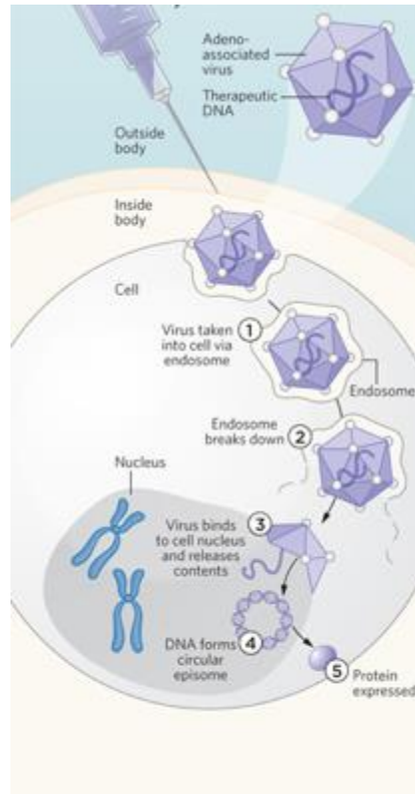
Integrating (Lenti)



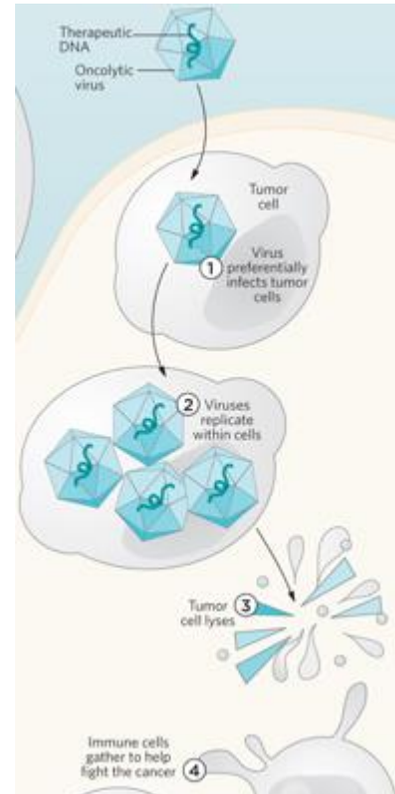
Ideal for *ex vivo* delivery to stem cell targets

Introducing Genetic Material via Viruses

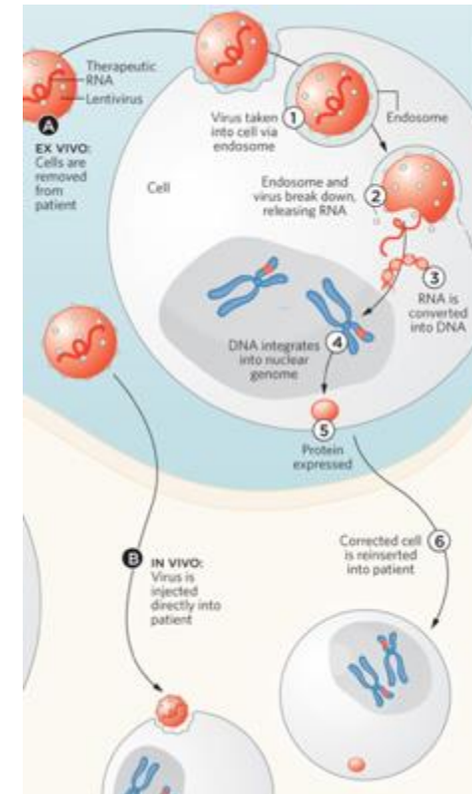
Non-Integrating



Integrating



Oncolytic



- Capsid – the protein shell of a virus; essential component involved in cell binding, internalization, and trafficking within the targeted cell
- Genes – the building blocks of inheritance
- Genetic disorder – results when genes don't produce the right proteins or don't produce them correctly
- Transgene – the gene or genetic material that is being transferred to the cell
- Vector – delivery vehicles that encapsulate therapeutic genes for delivery to the cell; include genetically disabled viruses, such as adeno-associated virus

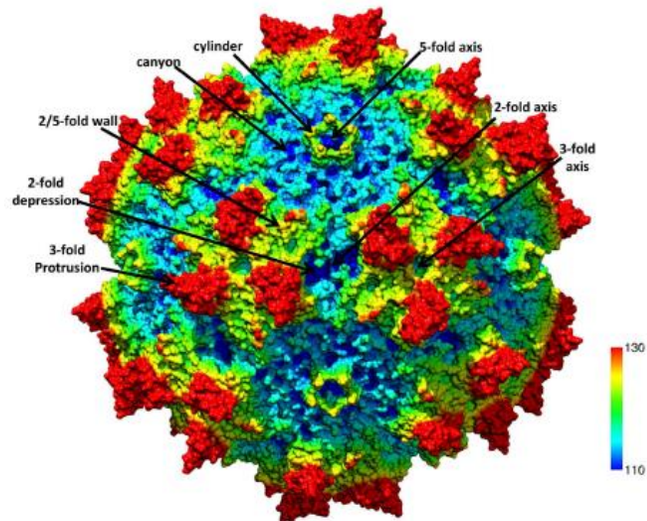
Why AAV Gene Therapy?

- Non-integrating - reducing oncogenic potential
- Multiple capsid types allowing for tailored tropism
- Expertise and experience to manufacture commercial scale product
- Leverages biopharma protein manufacturing experience

AAV: Family Parvoviridae, Genus Dependovirus

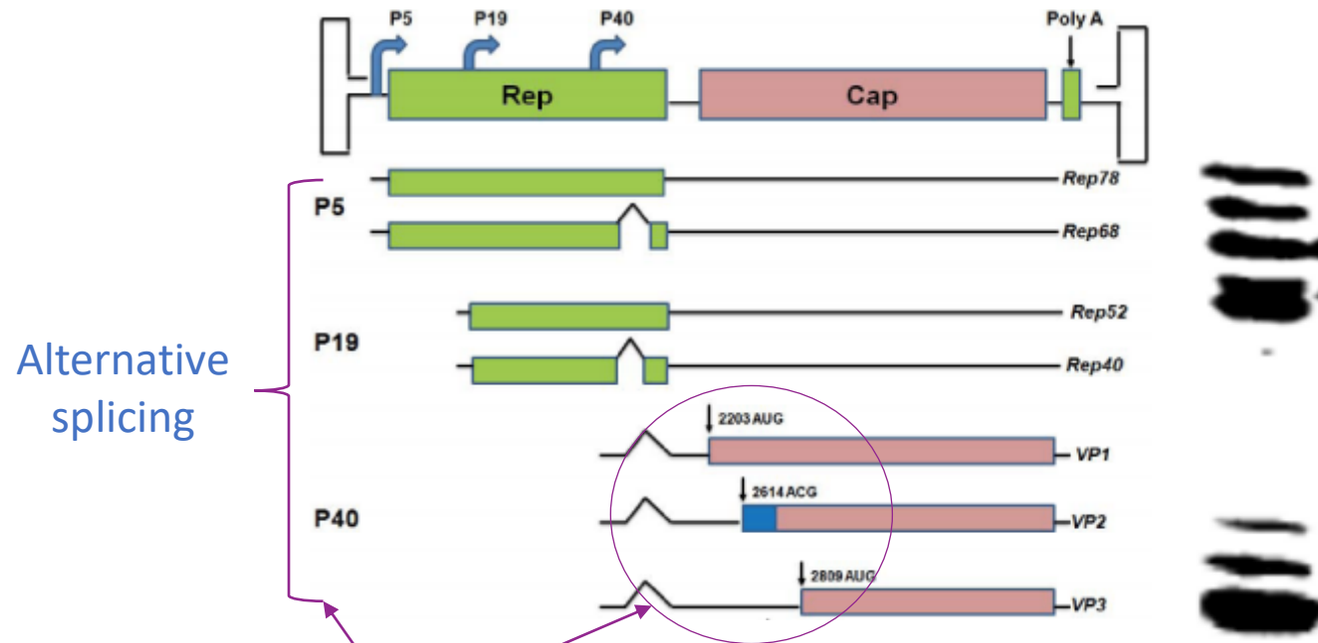
Scientific Platform

- AAV discovered in early 1960s
- Wild type AAV is not associated with disease
- Variable seropositivity in human population depending on capsid serotype



- 20 nm non-enveloped icosahedral capsid
- Virion extremely stable
- Single-stranded genome of 4,680 nt
- Three capsid proteins (VP1,2,3)
- Multiple capsid structural variants available

Dependovirus Genome Structure – Implications for Manufacturing

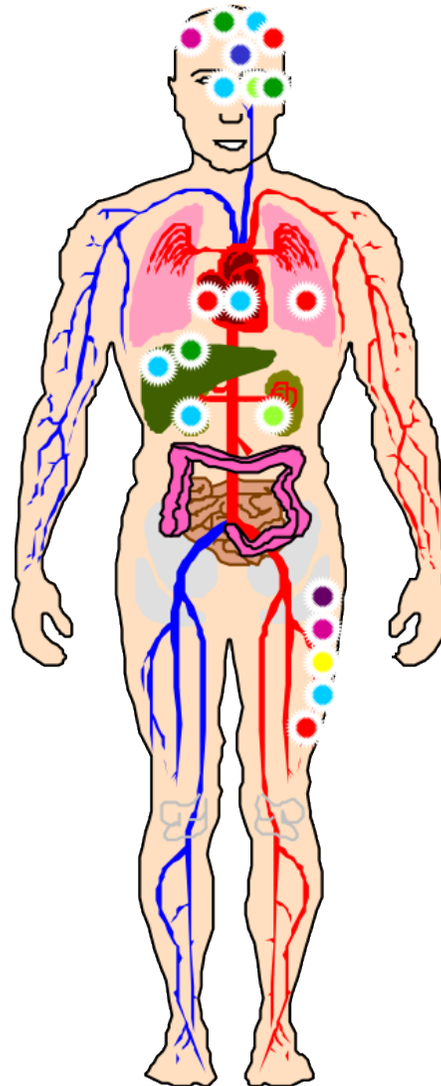
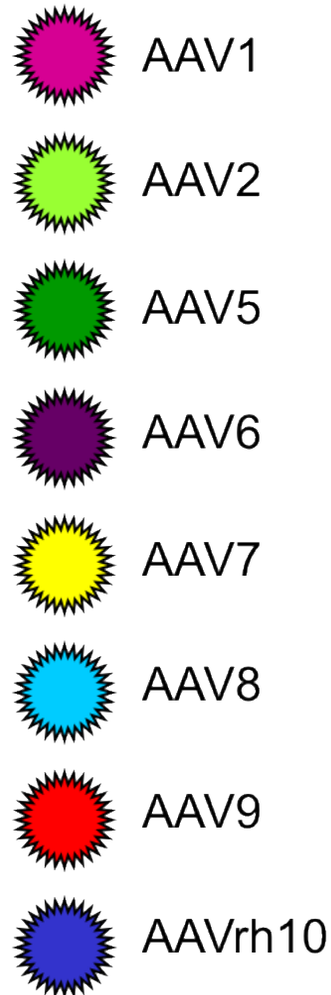


Alternative
splicing

And alternative
translation initiation

- Mammalian platform leverages existing molecular biology
- Baculovirus platform needs to engineer around this biology

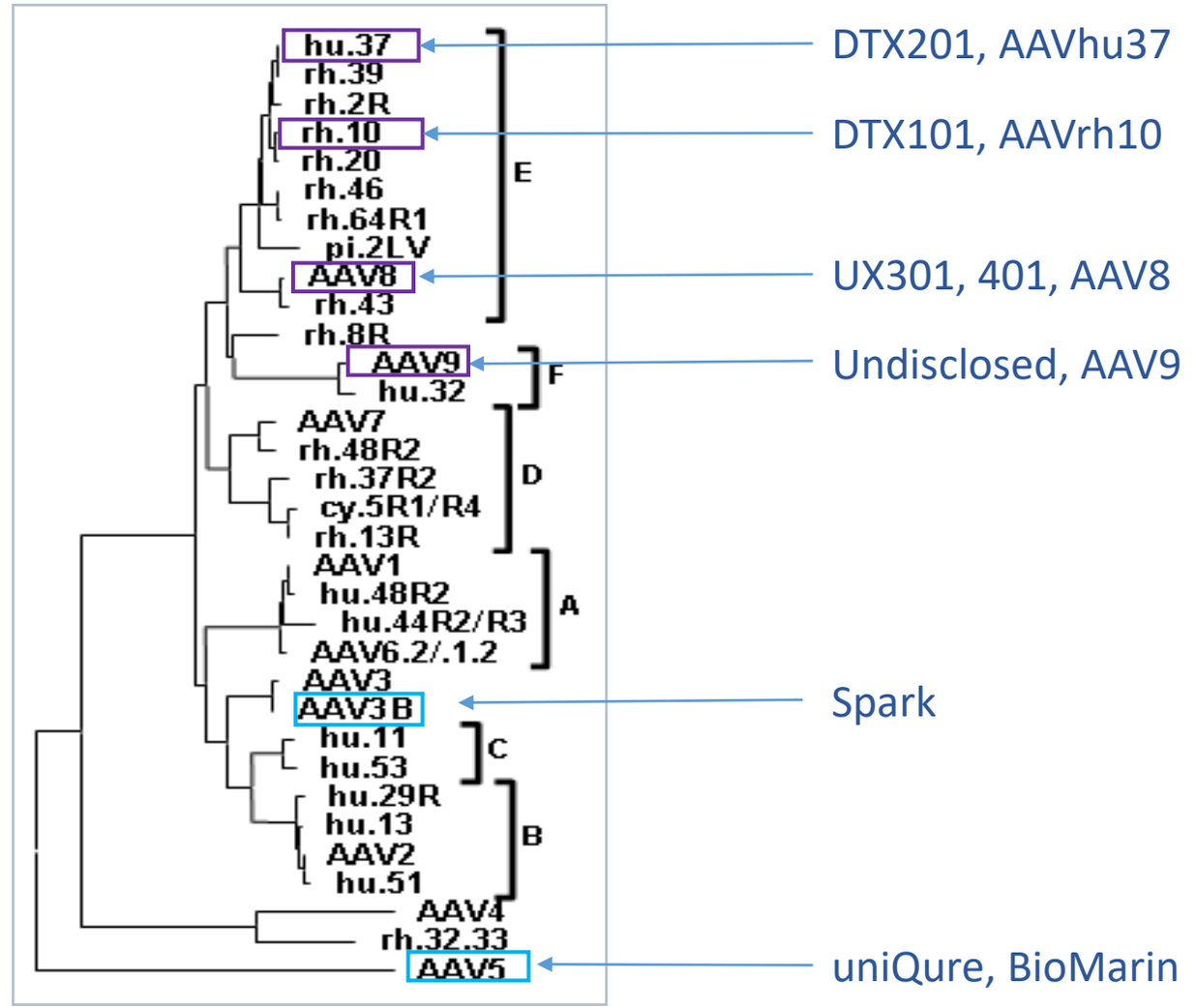
Different AAV Capsids Have Different Tropisms



- Tropism is not absolute
- Route of administration can overcome inherent tropism
- Prevalence of anti-capsid antibodies is a major consideration
 - Lack of precision on differences between capsids
 - Extremes are AAV2 with high antibody prevalence and AAV5 with low antibody prevalence

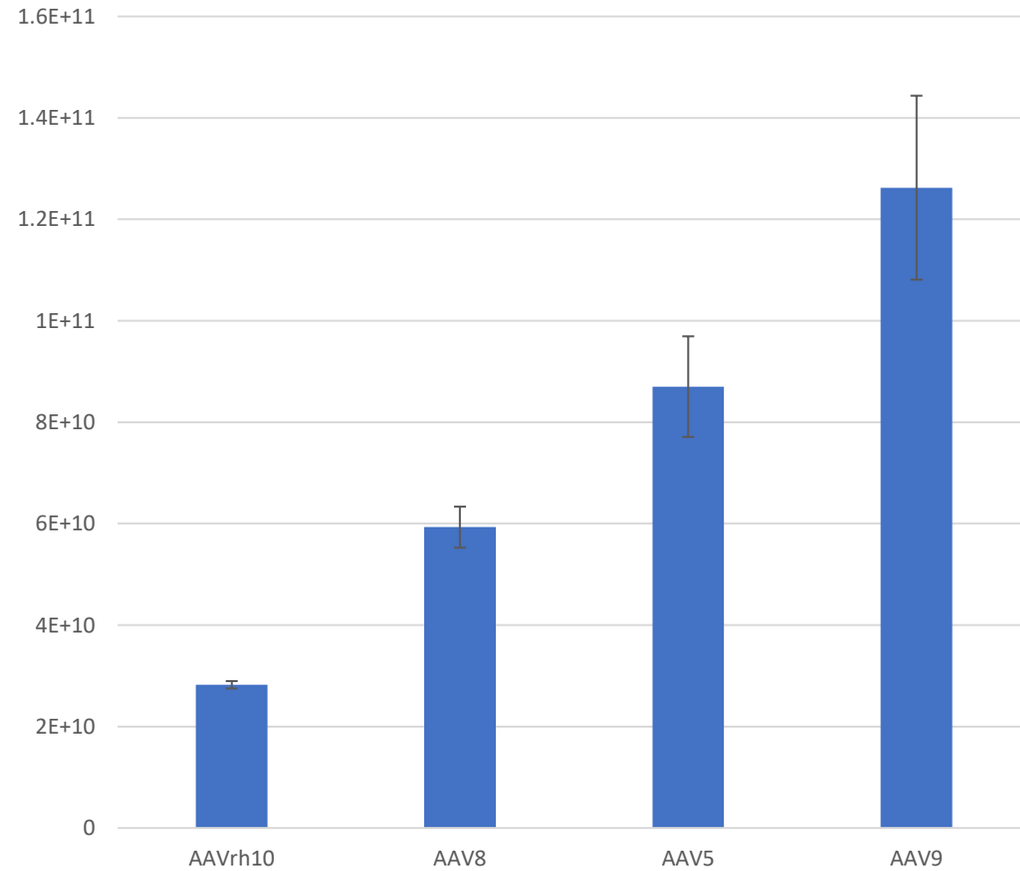
Current Products Based on Clade E Family

Capsids



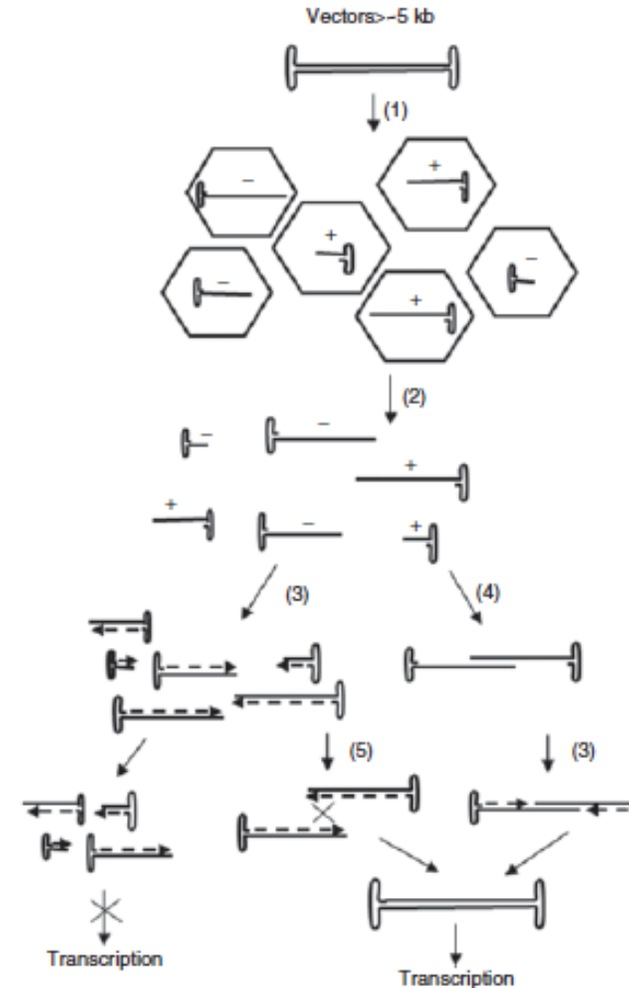
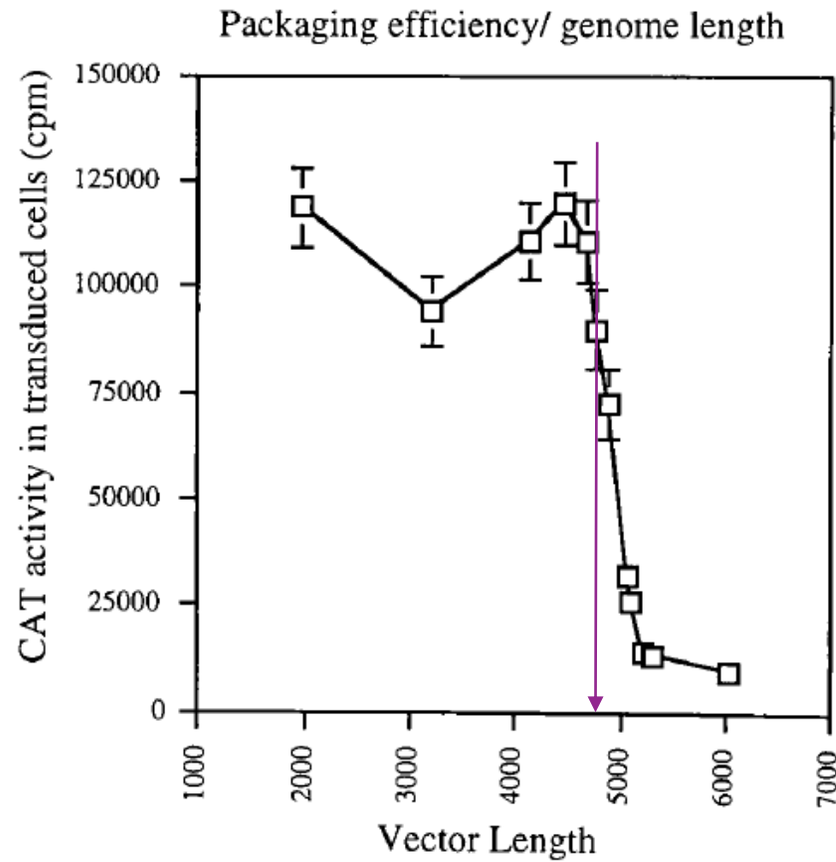
Ultragenyx Programs

Different Capsids May Have Manufacturing Advantages



- Packaging of 4 different vector genomes evaluated in 4 different capsids
- Certain downstream unit operations are common in vector purification, others are related but are fine-tuned

AAV Genome Forms – Reduced Packaging Efficiency & Expression Efficiency for Oversized Genomes



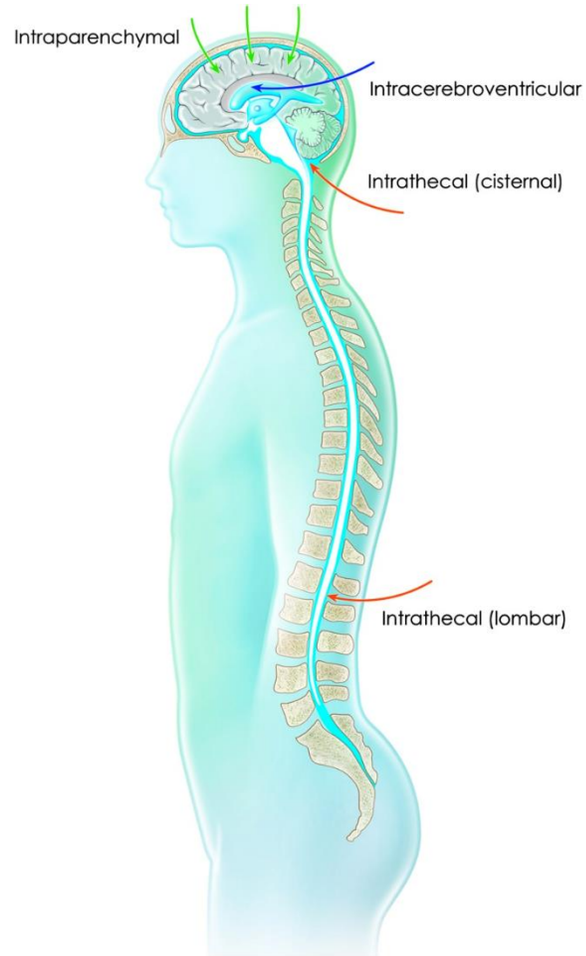
CNS AAV Gene Therapy

A number of viral vector-mediated phase I/II clinical trials have been initiated to treat neurologic disorders

Table 2
Viral vector-mediated clinical trials for neurological disorders.

| Disease | Vector | Transgene | Phase | Trial code |
|--|----------------------|---------------|-----------|---------------------------------------|
| Ex vivo | | | | |
| Alzheimer's disease | Retrovirus | NGF | I | US-0322 |
| Metachromatic leukodystrophy | Lentivirus | ARSA | I, II | Biffi et al., 2013 |
| Multiple sclerosis | Retrovirus | MBP | I, II | US-0851 |
| Wiskott-Aldrich syndrome | Lentivirus | WASP | I, II | Aiuti et al., 2013 |
| X-linked adrenoleukodystrophy | Lentivirus | ABCD1 | I, II | Cartier et al., 2009 |
| In vivo | | | | |
| AADC deficiency | AAV | AADC | I, II | NCT01395641 |
| Alzheimer's disease | AAV | NGF | I, II | NCT00087789, NCT00876863 |
| Batten disease | AAV | CLN2 | I | NCT00151216 |
| Batten disease | AAV | CLN2 | I, II | NCT01414985 |
| Canavan disease | AAV | ASPA | I | Leone et al., 2012 |
| Giant axonal neuropathy | AAV | GAN | I | NCT02362438 |
| Glioblastoma | Oncolytic poliovirus | – | I | NCT01491893 |
| Glioblastoma multiforme (GBM), other gliomas | Oncolytic adenovirus | – | I | NCT00805376, NCT01956734, NCT02197169 |
| Glioblastoma multiforme, other gliomas | Retrovirus | CD | I, II/III | NCT01470794, NCT02414165 |
| Glioblastoma, other gliomas | Oncolytic HSV | – | I | NCT02031965 |
| Glioblastoma, other gliomas | Oncolytic HSV | – | I | NCT00028158, NCT00157703 |
| Leber's hereditary optic neuropathy | AAV | MT-ND4 | I | NCT02161380 |
| Metachromatic leukodystrophy | AAV | ARSA | I, II | NCT01801709 |
| MPS IIIA (Sanfilippo Disease Type A) | AAV | SGSH, SUMF1 | I, II | NCT01474343, NCT02053064 |
| Parkinson's disease | AAV | GAD | I, II | NCT00195143, NCT00643890 |
| Parkinson's disease | AAV | NTRN | I, II | NCT00252850, NCT00400634 |
| Parkinson's disease | Lentivirus | TH, AADC, CH1 | I, II | NCT00627588, NCT01856439 |
| Parkinson's disease | AAV | GDNF | I | NCT01621581 |
| Parkinson's disease | AAV | AADC | I, II | NCT02418598 |
| Parkinson's disease | AAV | AADC | I | NCT00229736 |
| Pompe disease | AAV | GAA | I, II | NCT00976352 |
| Pompe disease | AAV | GAA | I | NCT02240407 |
| Spinal muscular atrophy type 1 | AAV | SMN | I | NCT02122952 |

Potential delivery sites for CNS AAV gene therapy



- **Global delivery**

- IV: easiest delivery, requires high vector doses, and may not target sufficient cells in regions of interest due to low penetration of BBB

- **CSF based delivery**

- ICV, IT (cisternal or lumbar): potentially challenging delivery method, may not reach deep brain structures, but will target a higher % of neurons compared to IV at a similar dose

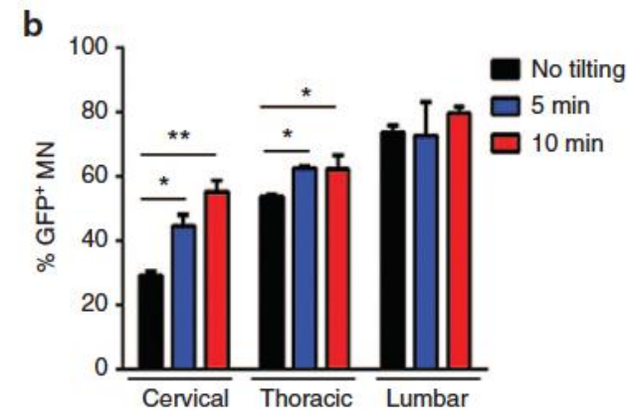
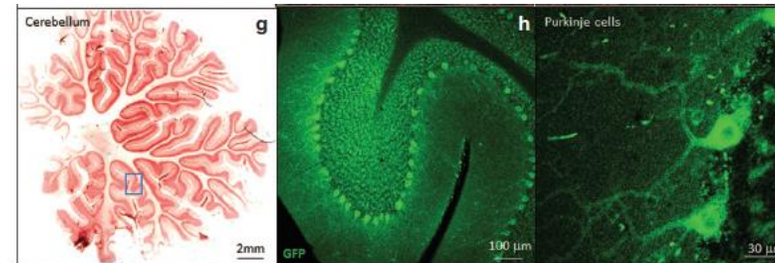
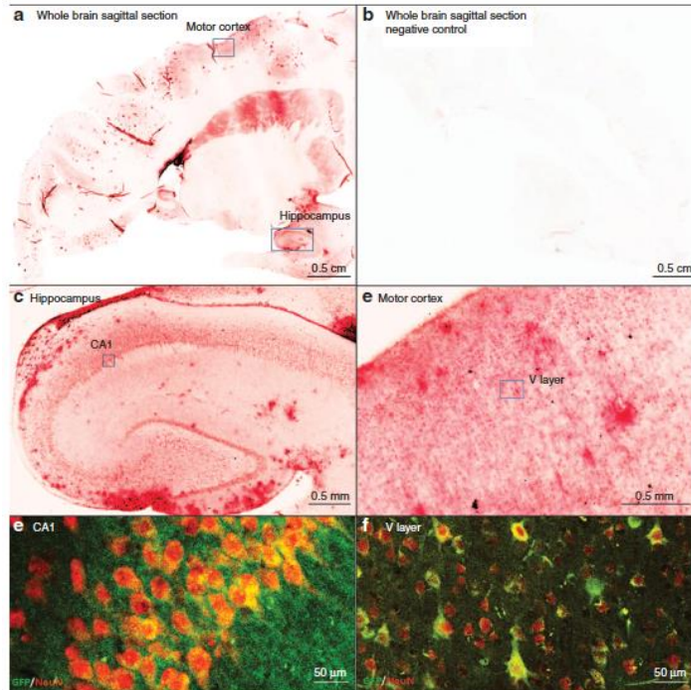
- **Intraparenchymal delivery**

- Potentially target a large percentage of neurons but only in a select area

IT delivery of AAV9-GFP results in expression throughout the brain

Meyer et al., 2015. *Molecular Therapy* (Brian Kaspar's lab)

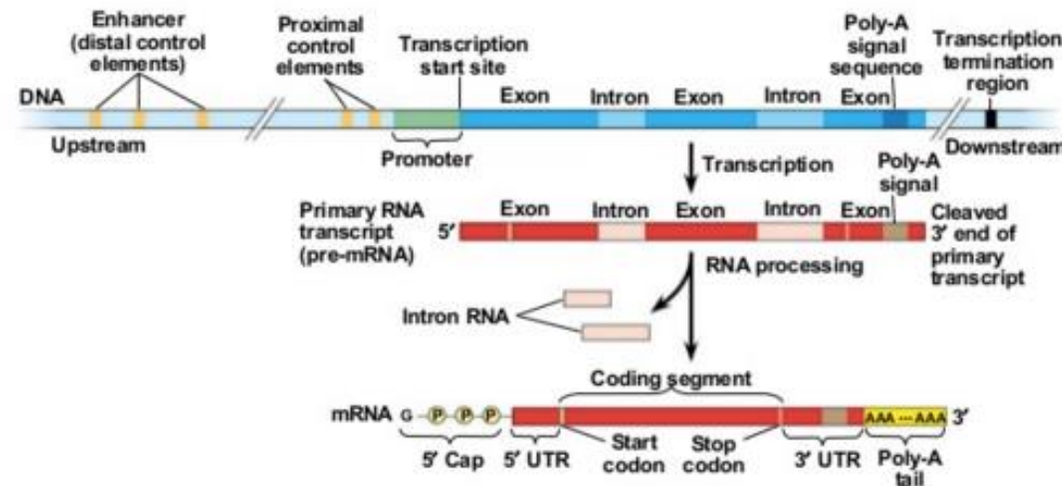
- scAAV9-CBA-GFP was delivered to n=5, **1 yr old NHPs (cynos)** via sacral-IT (**1×10^{13} vg/kg**) – used Trendelenburg position (head tilted down by 15-30 degrees for 10 min following infusion)
- Animals were sacrificed after 2 weeks
- **GFP was noted in all regions of the brain, with particularly strong signal in the hippocampus, motor cortex and cerebellum**



Note: others have reported similar results but with less robust brain delivery

DNA Component: Molecular Engineering to Squeeze Genes into the AAV Genome Limit (5 -5.4 kb)

- Most therapeutic gene coding sequences will use their minimal cDNA format
- Overstuffing capsid has negative yield and quality consequences
- Additional elements, such as introns & 5' and 3' UTR sequences, can sometimes be included
- Greatest technical challenge is engineering small enhancer and promoter combinations to achieve tissue specific gene expression

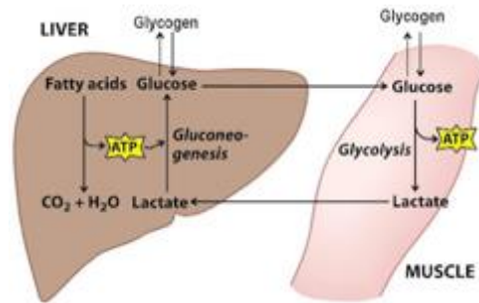
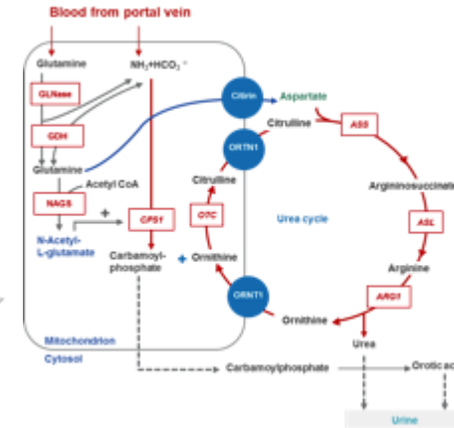


Targeting the Liver – A Master Hub for Rare Diseases

Hemophilia

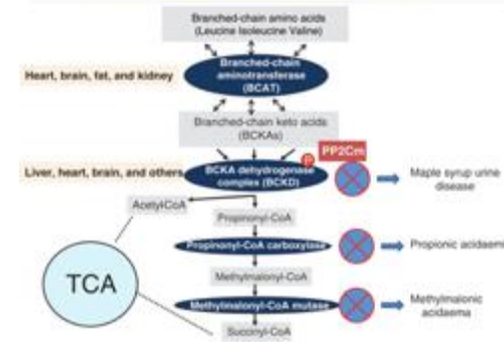


Urea Cycle Disorders



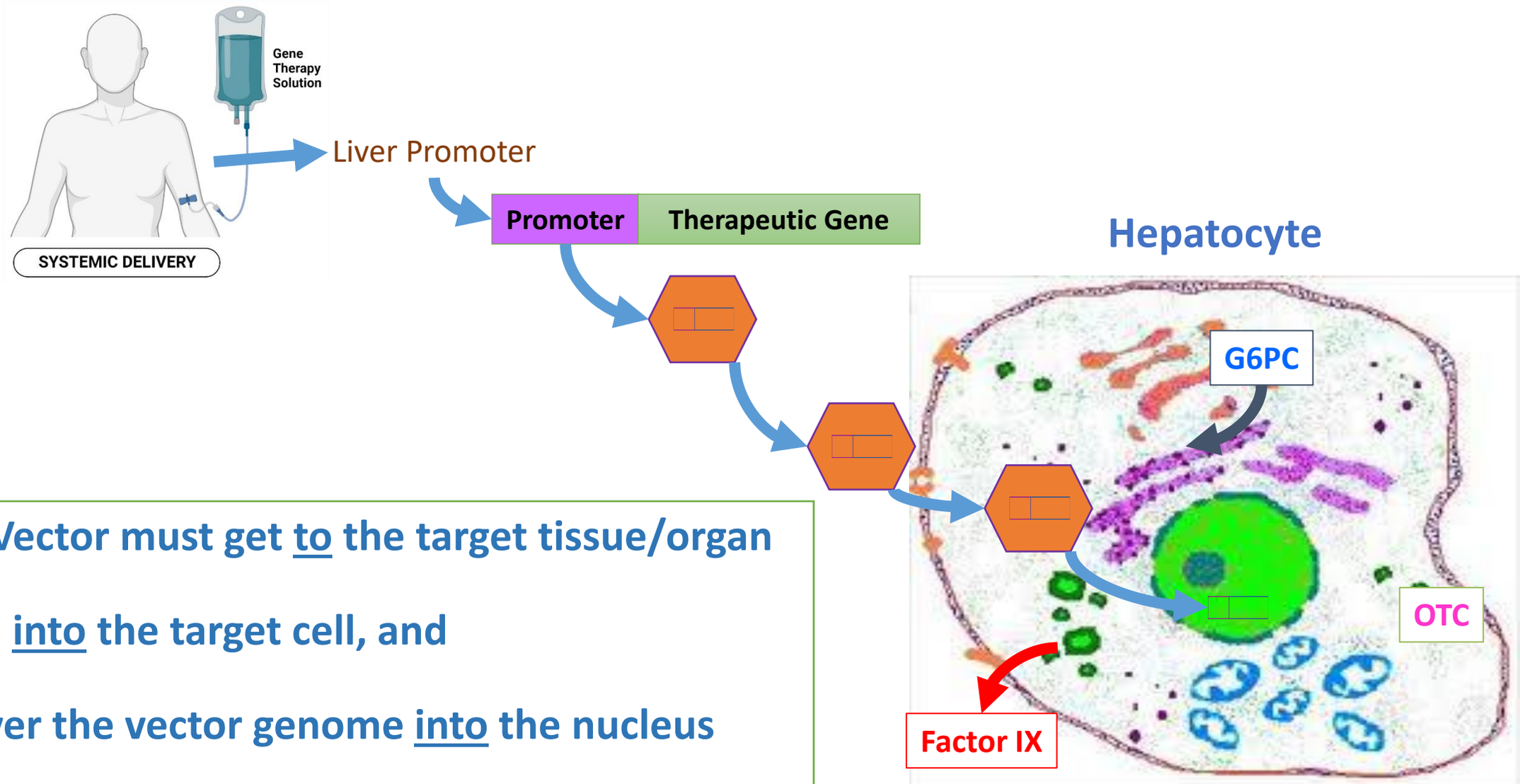
Storage Disorders

Catabolism of branched-chain amino acids



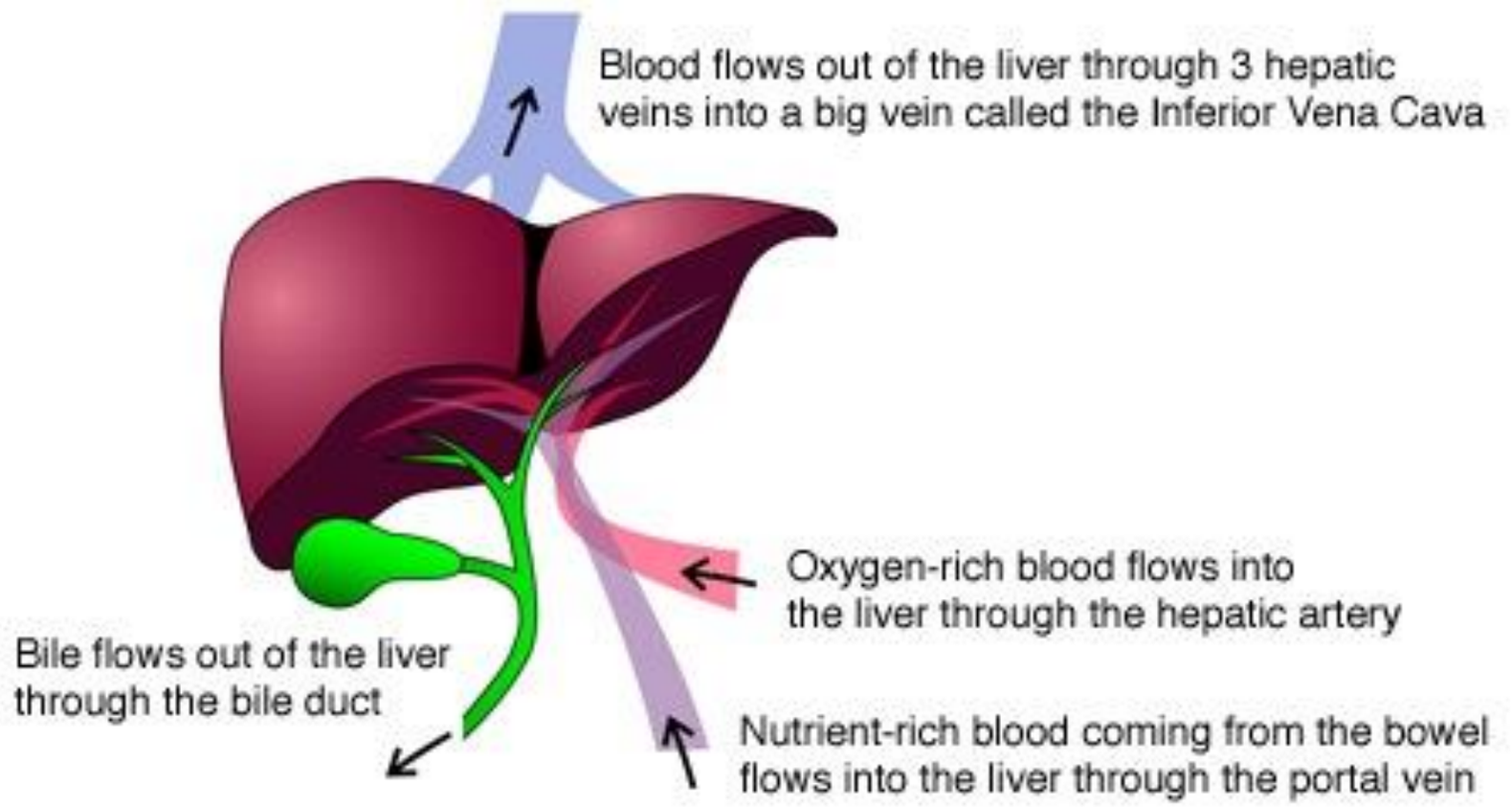
Aminoacidopathies

Gene Therapy Delivery is a Complicated, Multi-Step Process

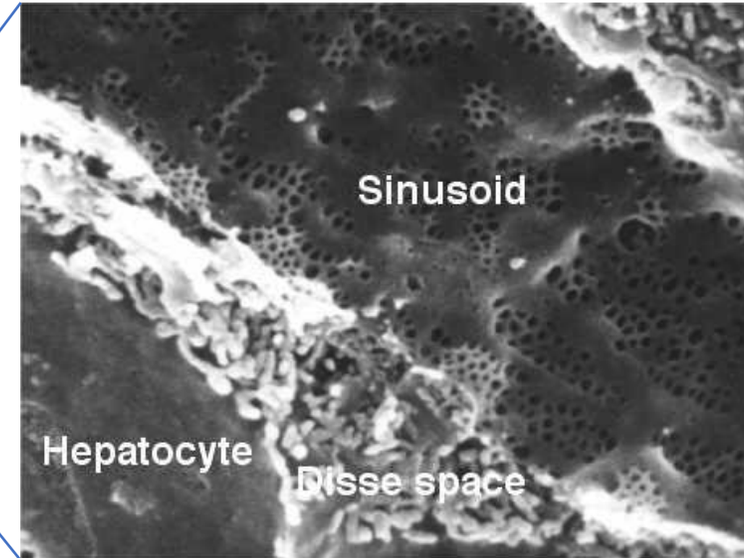
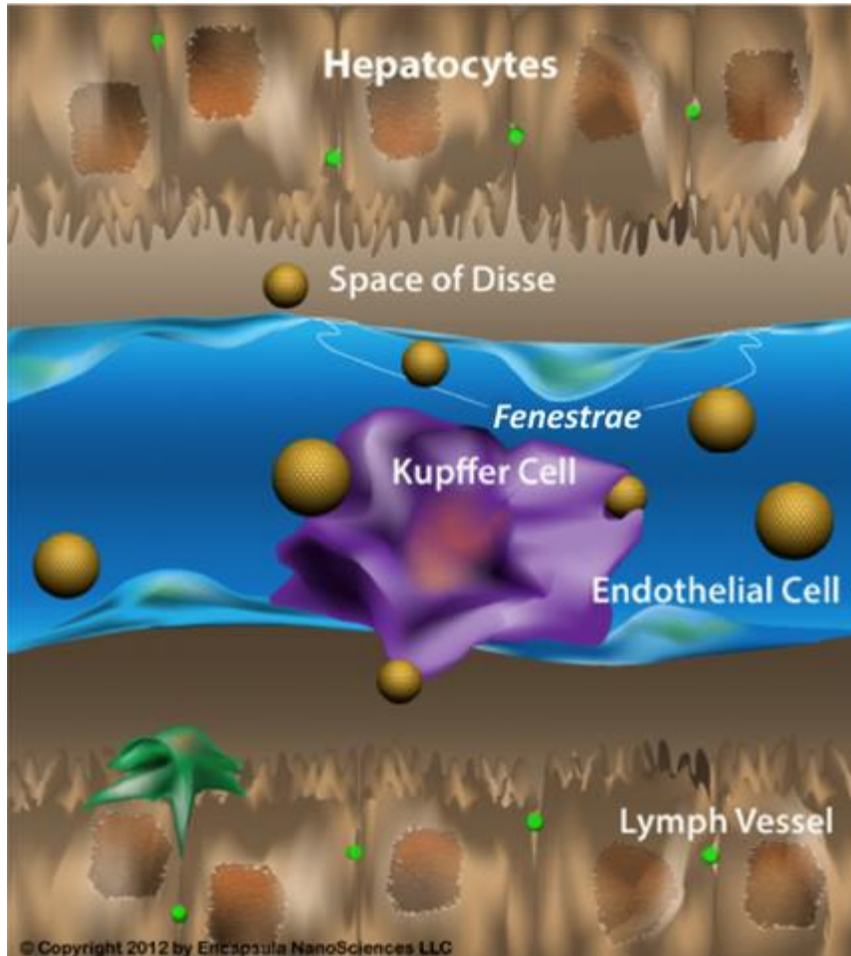


1. The Vector must get to the target tissue/organ
2. Then into the target cell, and
3. Deliver the vector genome into the nucleus
4. Where the therapeutic gene must be expressed

Blood Flows Through the Liver at 1.5L per min



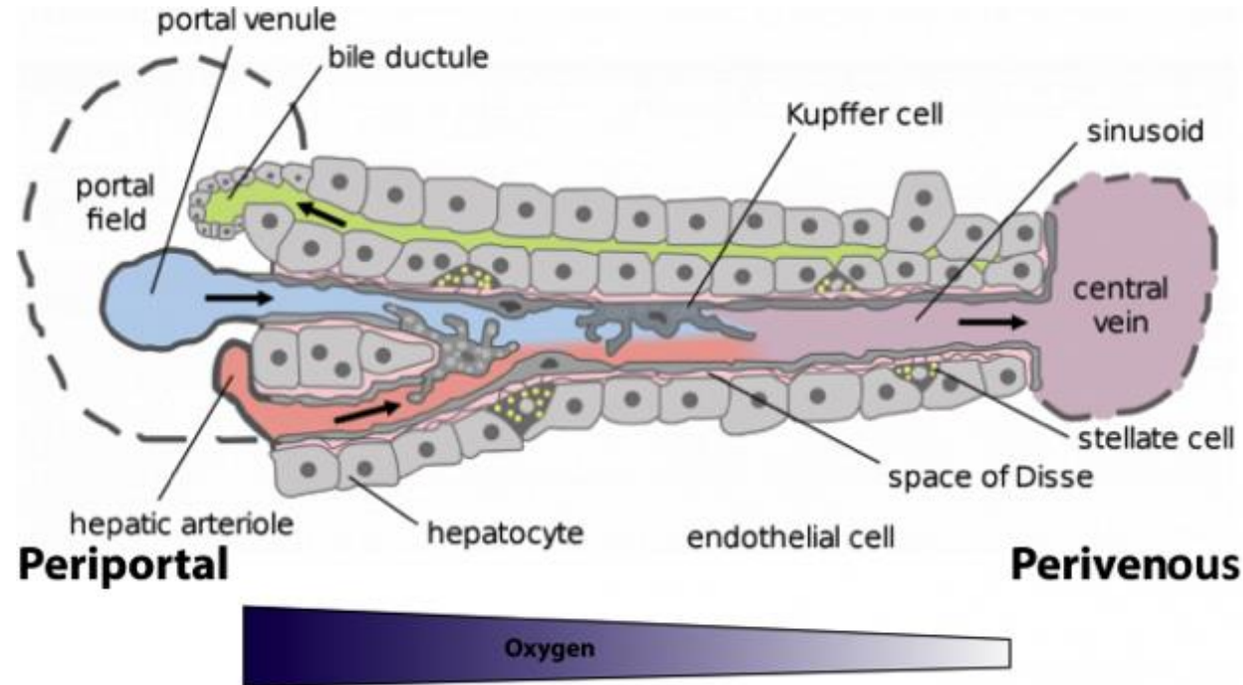
Liver Sinusoidal Endothelium Is Fenestrated



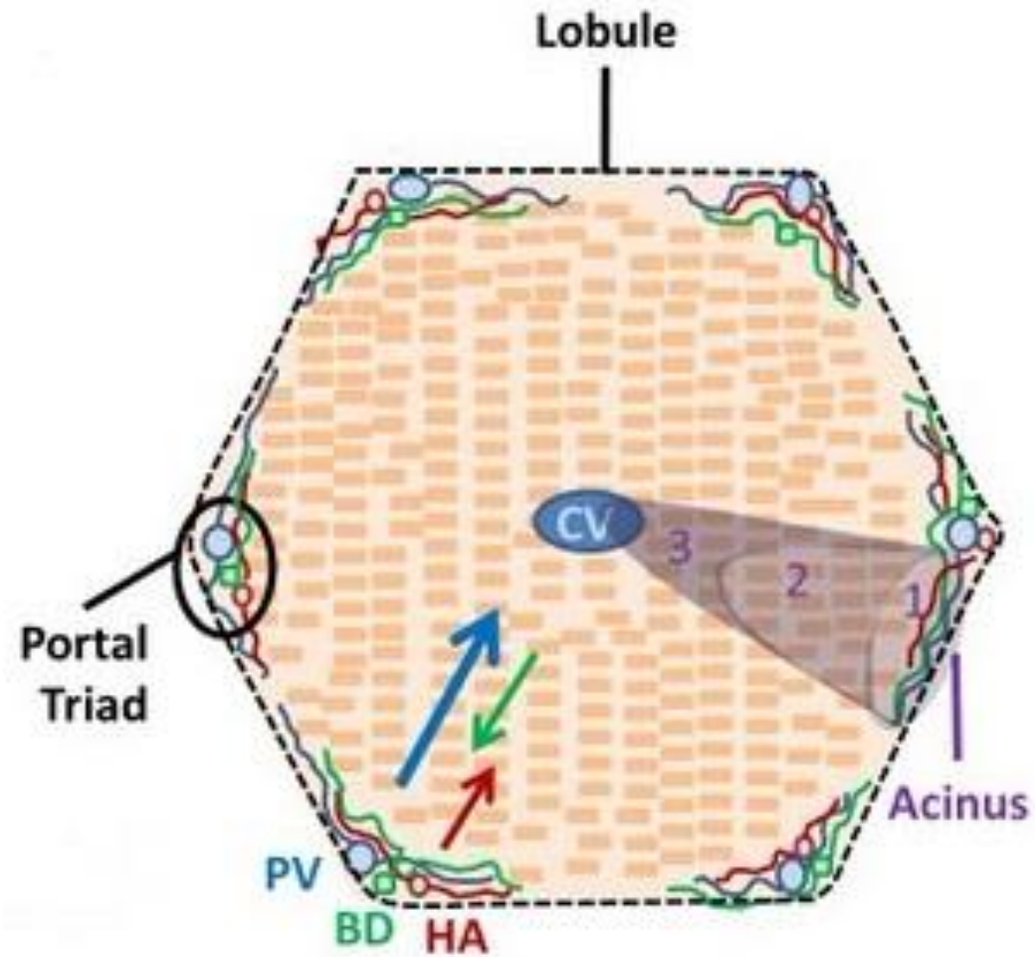
Hepatocytes Are Divided Into Functional Zones

High O₂ processes

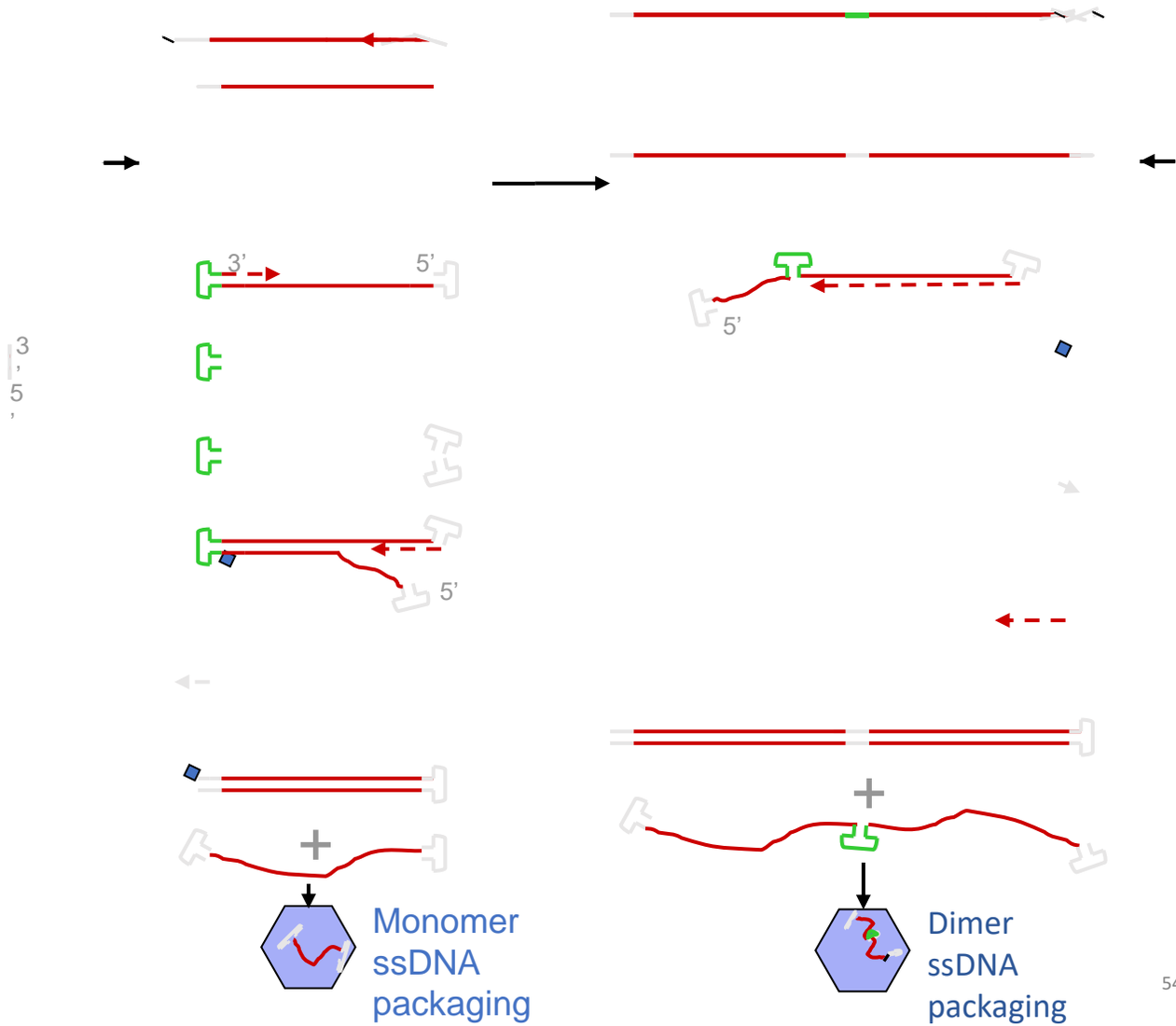
Low O₂ processes



Functional Zones of the Liver (the big picture)

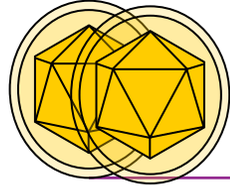


AAV Genome Forms – Single Strand versus Self Complementary



AAV Vector Manufacturing – HSV Advantages

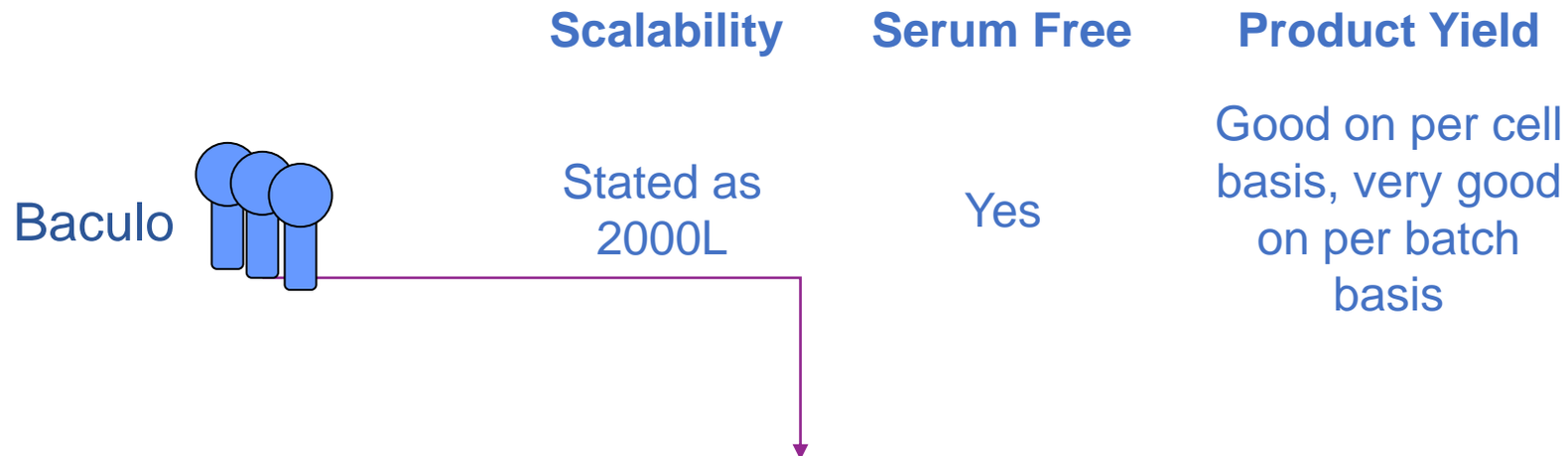
and Limitations

| | Scalability | Serum Free | Product Yield |
|---|---|------------|---|
| HSV  | Bioreactor format enabled, scale unknown (100 to 200L ??) | Yes/No | Good on per cell basis, moderate on per batch basis |

- Comparability to other mammalian platforms good
- Key features/bugs – Good product quality, easy removal of rHSV/Limited by stability of rHSV, requires 2 rHSV vectors as complex GMP raw material supply train
- Alternate bioreactor format would be required to implement at Woburn; different format for vector engineering

AAV Vector Manufacturing – Baculo

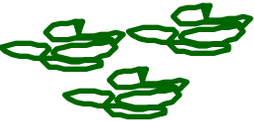
Advantages and Limitations



- Comparability to other mammalian platforms a question
- Key features/bugs – high yield & serum free, high cell density/requires 2 or 3 rBac viruses as supply chain, non-optimal stability of rBac (genetic & storage), complex molecular engineering required to support fidelity of AAV life cycle
- EMA Glybera Assessment highlights several gaps in CMC
- Alternate bioreactor format would be required to implement at Woburn; different format for vector engineering

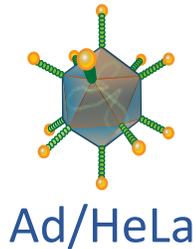
AAV Vector Manufacturing – HEK293

Advantages and Limitations

| | | Scalability | Serum Free | Product Yield |
|--------------------|---|--|------------|---|
| Plasmid Adherent |  | Poor | No | Good on per cell basis, low on per batch basis |
| Plasmid Suspension |  | Data to 200L, probably could go higher | Yes | Good on per cell basis, good on per batch basis |

- Comparability to adherent 293 high
- Key features/bugs – HEK293 platform used for majority of AAV gene therapy trials/plasmid represents high GMP raw material burden
- Readily implemented at Woburn; leverage our core AAV science, support basic PD, de-risk CMO tech transfer

and Limitations



Scalability

Proven to
2000L
Bioreactor

Serum Free

Yes

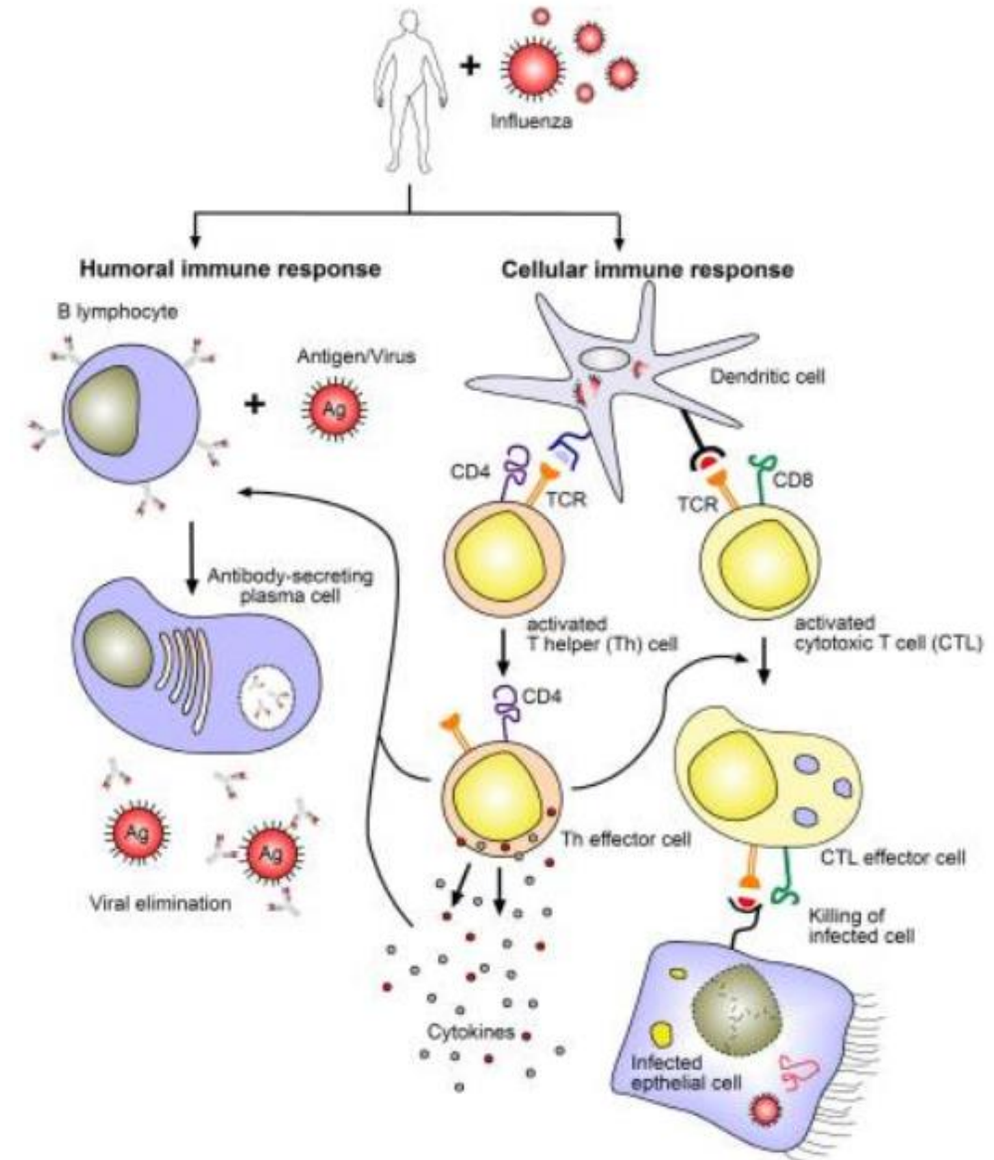
Product Yield

Very good on per
cell basis, very
good on per
batch basis

- Comparability to other mammalian platforms good
- Key features/bugs – only clonal system enabling screen for desired features; E/F, rcAAV, other requires high titer Ad helper GMP raw material supply train
- Readily implemented at Woburn; leverage from our core AAV science, support basic PD, de-risk CMO tech transfer

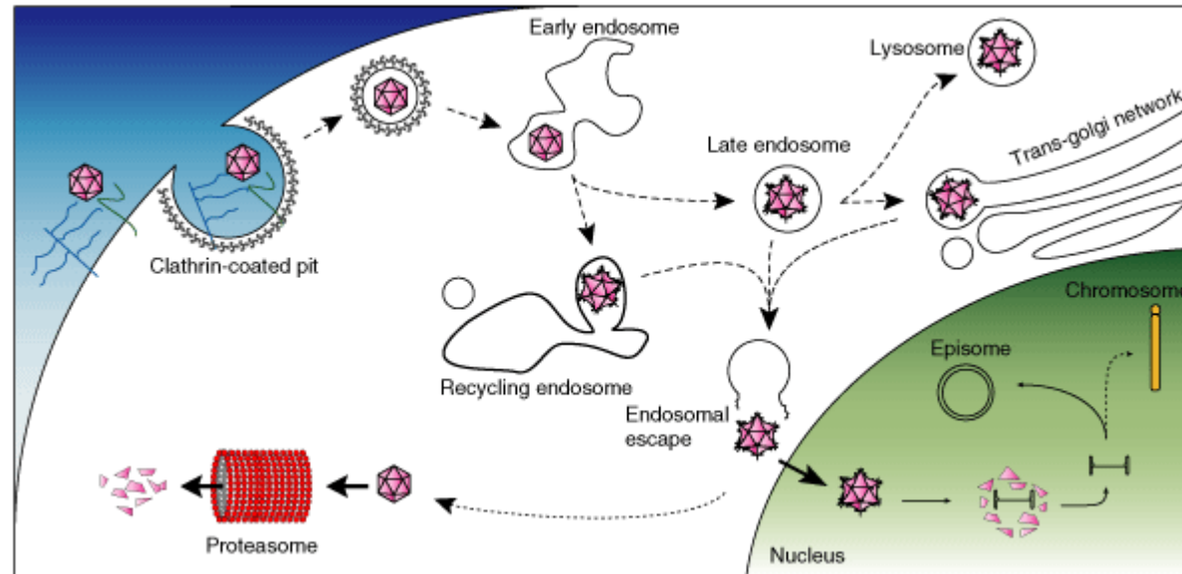
The Immune System – A Primer

- Two pathways – humoral and cellular
- Key signaling includes T-cell, B-cells and NK cells
- AAV vectors retain some viral coding that may be recognized as foreign
 - Capsid
 - Transgene
 - Genome



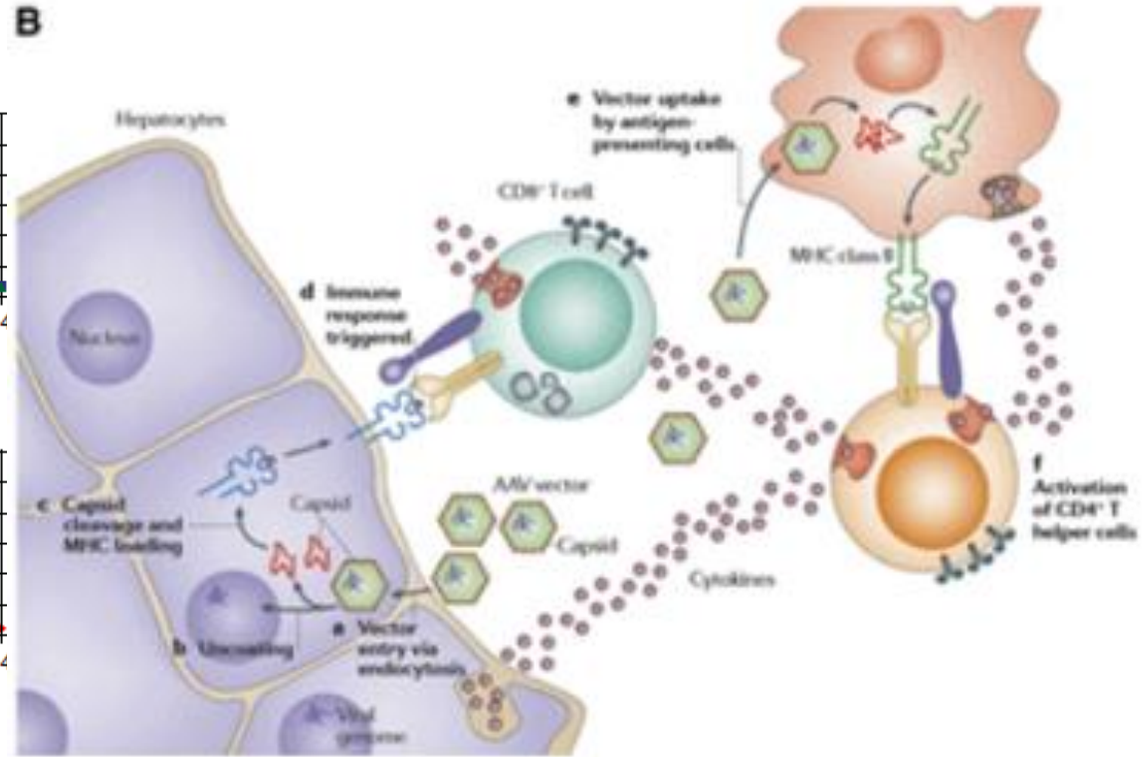
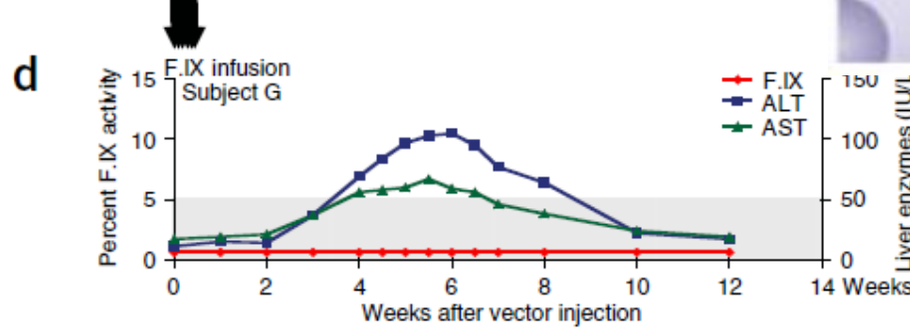
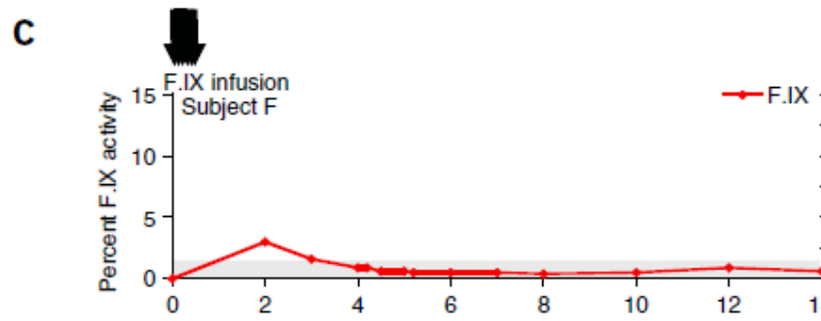
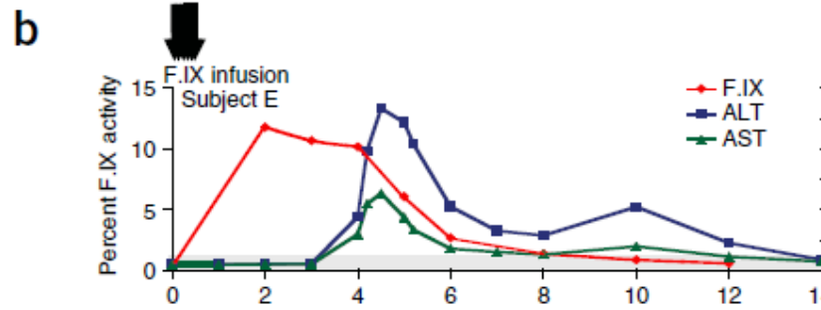
Potential Goals of immune intervention in AAV GT

- T cell responses to capsid antigens?
- T cell responses to therapeutic protein?
- B cell responses to capsid antigens?
- B cell responses to therapeutic protein?



First Observation of a Reduction in AAV Gene

Therapy – First Liver Delivered Hemophilia Trial



First Introduction of Steroids to Address Reduction in Efficacy – Nathwani 2011

- Steroids broadly suppress the immune system
- Nathwani's use of steroid was likely serendipitous
- Outcome was favorable and patients have sustained activity beyond 6 years

