GGSB Prelim Q2 – Hang Chen

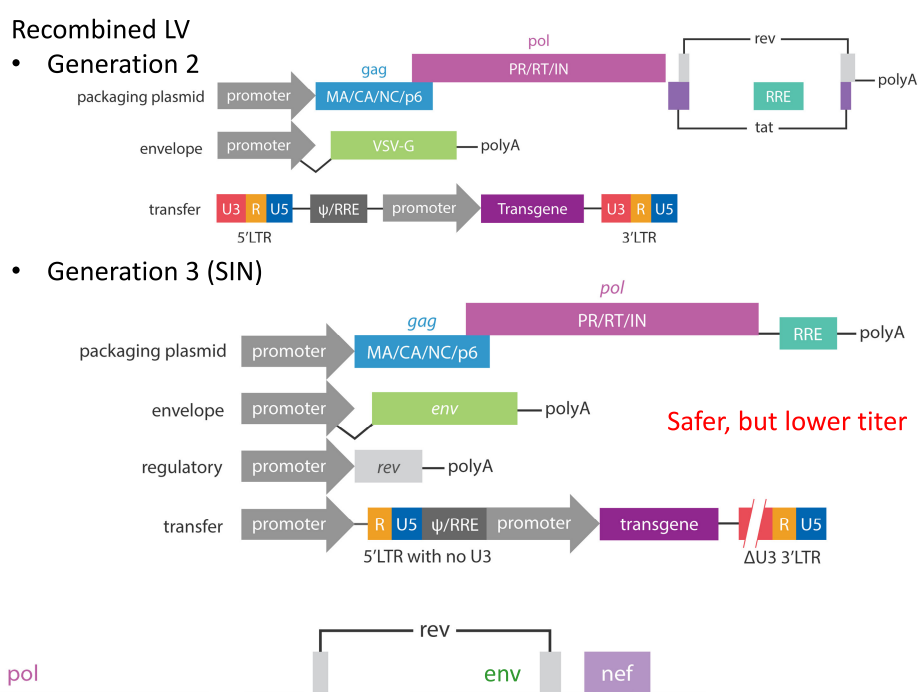
Gene therapy: treating genetic disorders using genetic approaches

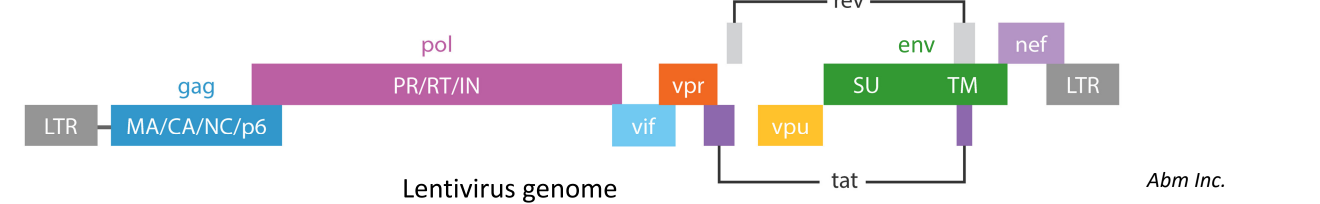
- Major class of diseases that are conducive to gene therapy:
 - Monogeneic: easy to target
 - Loss-of-function or haploinsufficiency: easy to rescue
 - Many are also hematological diseases: easy to access
 - Lower-level restoration: easier to molecularly fix cells than tissues

| Dx | Cause | Gene | Gene changes | Location | Vector |
|---|--|------------------------|--|----------|--------|
| X-linked Severe Combined Immunodeficiency (SCID) | Lymphocytes development deficiency | IL2RG | Mutations in <i>IL2RG</i> gene | Ex vivo | RV |
| Thalassemia | Less alpha or beta hemoglobin | HBA1 and/or HBA2 | Mutations in <i>HBA1</i> and/or <i>HBA2</i> gene | Ex vivo | AAV |
| Sickle cell anemia | Hemoglobin polymerization | НВВ | A mutation in both HBB alleles | Ex vivo | AAV |
| Congenital hemophilia | Blood does not clot properly | F8 and/or F9 | Mutations in F8 or F9 genes | In vivo | AAV |
| Vision Loss | Impaired retinoid cycle | RPE65 | Mutations in <i>RPE65</i> gene | In vivo | AAV |
| Spinal muscular atrophy | Insufficient survival motor neuron protein (SMN) | SMN1 | Mutations in SMN1 gene | In vivo | AAV |

Lentivirus (LV)

- Type
 - Retrovirus (enveloped)
- Genome
 - Single stranded positive sense RNA
 - ~10kb
 - 5' and 3' LTRs: required for transcription and reverse transcription
 - gag: core
 - pol: reverse transcription
 - env: surface protein
 - rev/tat: regulatory
 - some other replication related genes





Recombined Lentivirus (LV)

- Infection
 - Tropism: VSV-G, for most cells
 - Entry: membrane fusion and endocytosis
 - Dividing cells (hard to cross nuclear membrane)
 - Genome integration

• Pros:

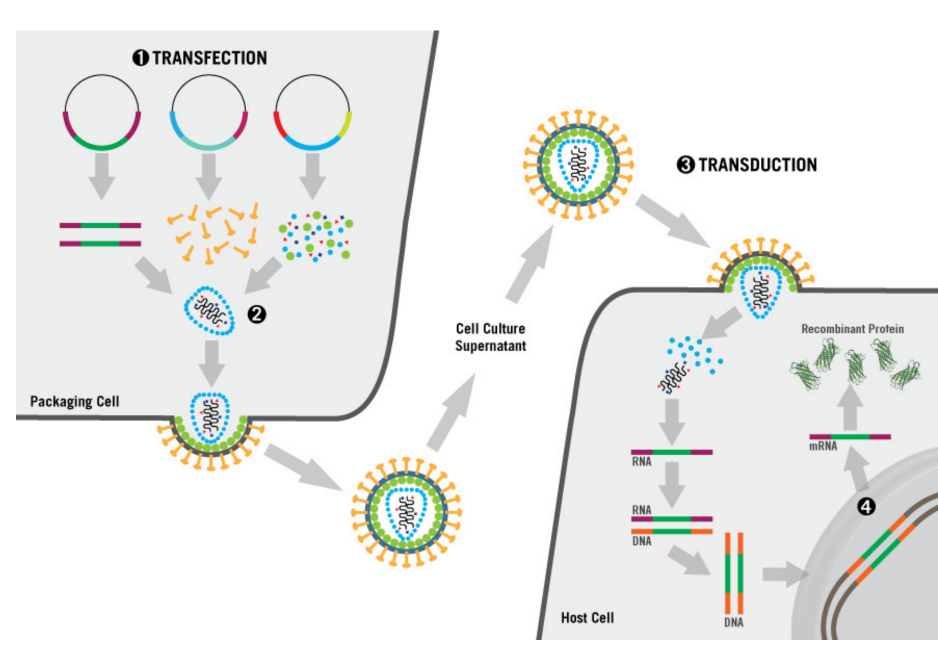
- Large capacity for transgene
- Easy to design (transfer plasmid)
- SIN system for safety
- Integrated into genome, permanent expression

• Cons:

- Insertional mutagenesis risks
- Copy number issues

Suited for:

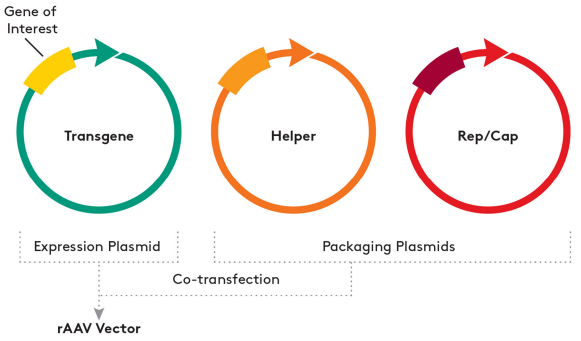
- Dividing cells
- Permanent correction
- E.g., hematopoiesis diseases (ex vivo treatment)

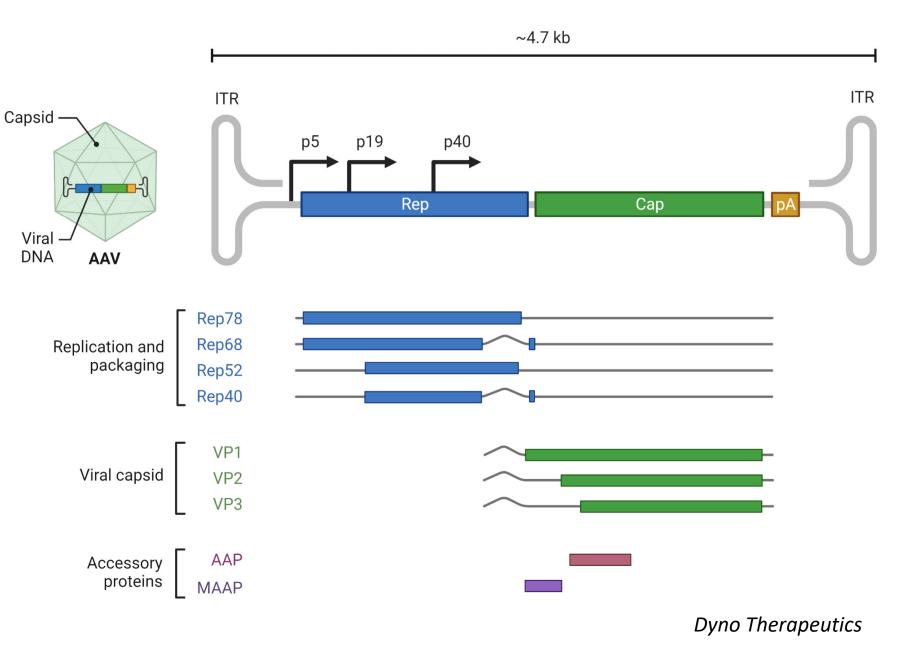


Mirus Bio LLC

Adeno-associated virus (AAV)

- Type
 - Dependovirus (non-enveloped, rely on other viruses to help replicate)
- Genome
 - ssDNA
 - ~4.7kb
 - 5' and 3' ITRs: required for the synthesis of the complementary DNA
 - rep: replication genes
 - cap: capsid genes





Recombined AAV (rAAV)

- transfer plasmid: ITR and transgene
- helper plasmid: required for virus replication
- rep/cap plasmid: may require special design for targeted cells

Recombined Adeno-associated virus (AAV)

- Infection
 - Tropism: more specific (may require optimization, e.g., directed evolution, in silico design)
 - Entry: endocytosis
 - Both dividing and non-dividing cells
 - Very low genome integration

• Pros:

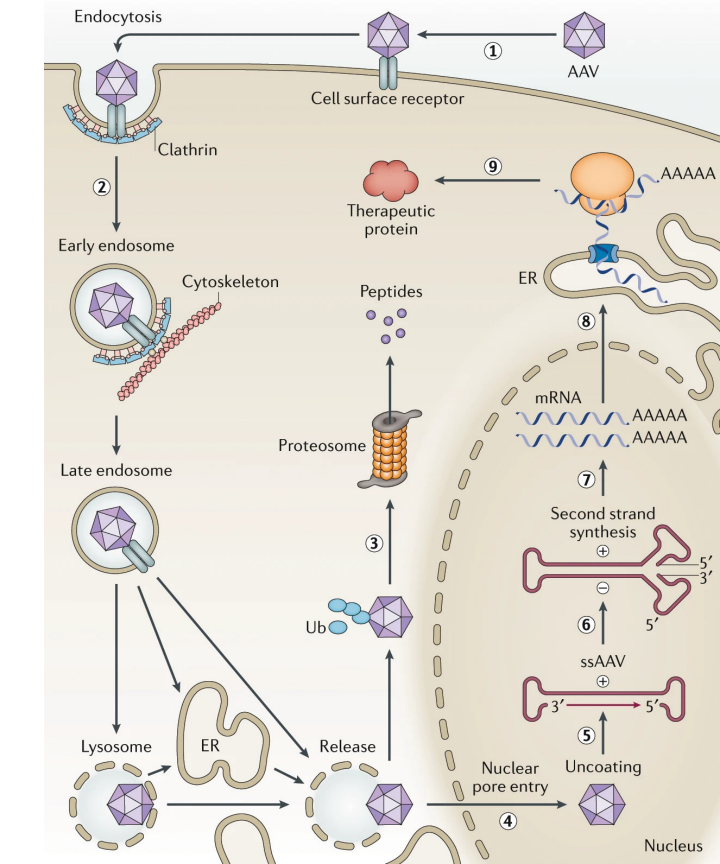
- Tissue specificity
- Very low insertional mutagenesis risks (although it could happen if administrated in long term)
- AAV itself does not cause disease

Cons:

- Small capacity for transgene
- Complicated to design, hard to manufacture in large scale
- Some people may have immunity against naturally existing AAV, which can decrease the efficacy

• Suited for:

- Non-dividing cells
- Transient expression
- E.g., neurological disorders (in vivo treatment)



Challenges

- Ethics
 - Bottomline: no germline editing
- Safety
 - Genotoxicity (insertional mutagenesis, copy number issues)
- Efficacy
 - Delivery (tropism, immunity, cell migration)
- Cost
 - Up to \$1.2 billion per patient
- Some other directions:
 - ASO (RNAi issue)
 - CAR-T CAR-NK (exhaustion issue)
 - Maybe SARS-CoV-2 as a vector in the future? (~30Kb capacity, strong infection ability)