



# Manufacturing Strategy for Rare Diseases

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# Agenda

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

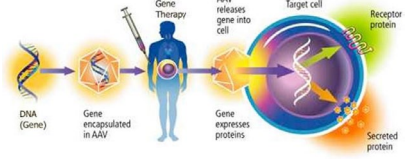
- Framing the Manufacturing Challenge
- How Ultragenyx Addresses These Challenges
- Manufacturing, QC and Risk across life cycle
- CMO Management, GxP requirements
- Q&A

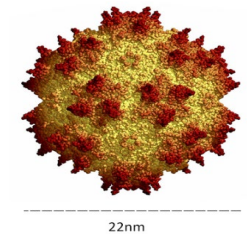
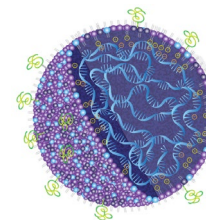
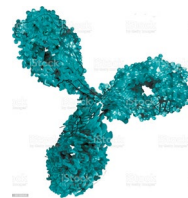
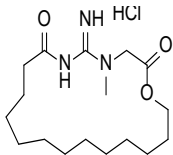
# Framing the Manufacturing Challenge

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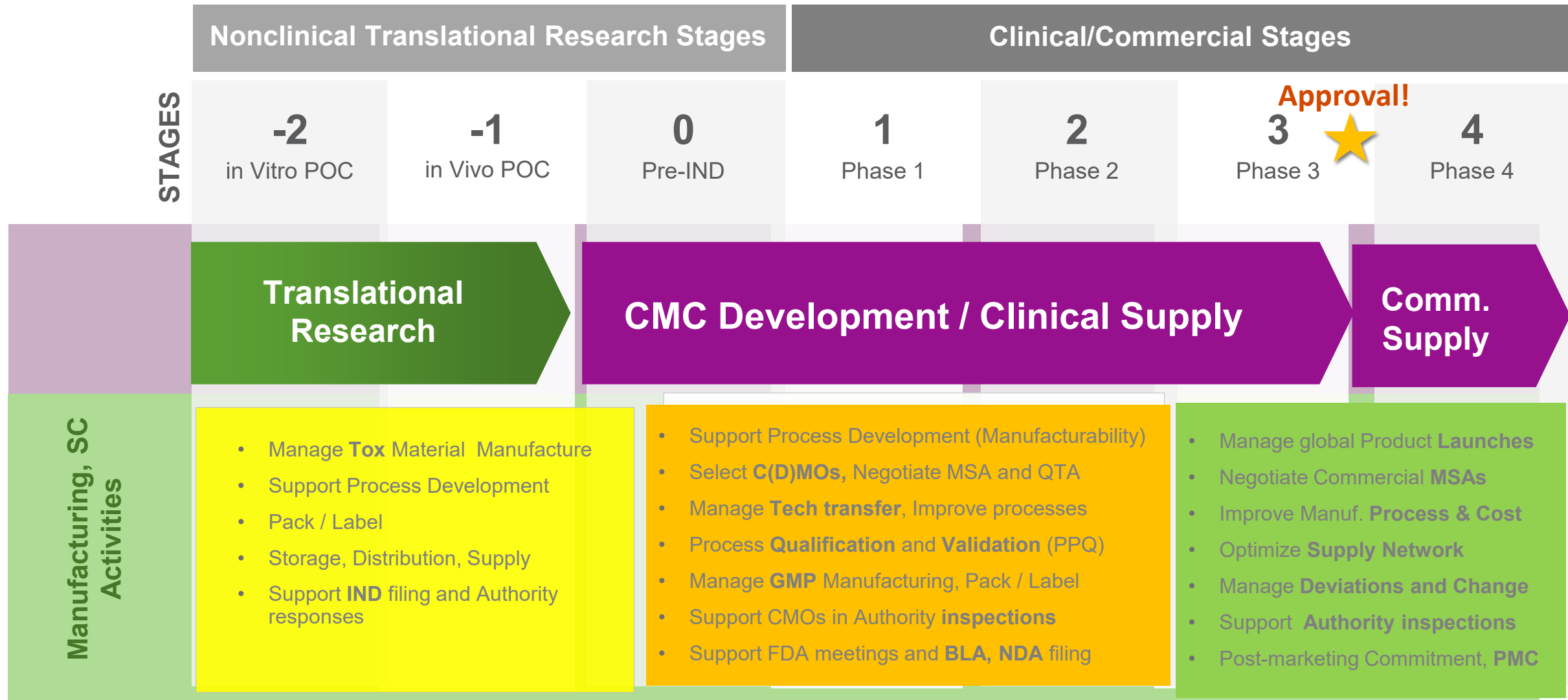
- Know the “race” you are running
  - eg; Fastest to IND, Fastest to Approval, Cost effective to IND, Cost efficient until value inflection point?
- What modality to use?
  - Biology is biology
  - Understand the tradeoffs with the different modalities
- Be as explicit about the risks as you can
  - What is sufficient?
  - What scale should I operate at?
  - Technical and scientific
  - Regulatory
  - Benefit / risk
- Recognize you are learning too

# Ultragenyx Pipeline by Modality and Stage

	Small Molecule	Traditional Biologics	Nucleic Acid Therapy	AAV Gene Therapy
Commercial	 <p><b>Dojolvi</b></p>	<p><b>Burosumab</b> <b>Mepsevii</b> <b>Evkeeza</b></p> 		
Clinical	<p><b>ATX95 (mRNA)</b> <b>Prednisolone (GT)</b> <b>Na Acetate (GT)</b></p>	<p><b>UX143</b></p> <p><b>Plasmid in E.coli</b> <b>(GT and mRNA)</b></p>	<p><b>GTX102 (ASO)</b></p> <p><b>UX053 (mRNA)</b></p>	<p><b>DTX301</b> <b>DTX401</b> <b>UX701</b> <b>UX111</b></p>
Pre-Clinical	<p><b>UX068</b> <b>UX016</b></p>	<p><b>UX100 (E.coli)</b></p>		<p><b>UX055</b> <b>UX810</b></p>



# Key Manufacturing Activities Along Lifecycle



# Testing also Has a Significant Lifecycle to Consider as Well

**Analytical comparability may need to occur during the lifecycle**

**Analytical Development**

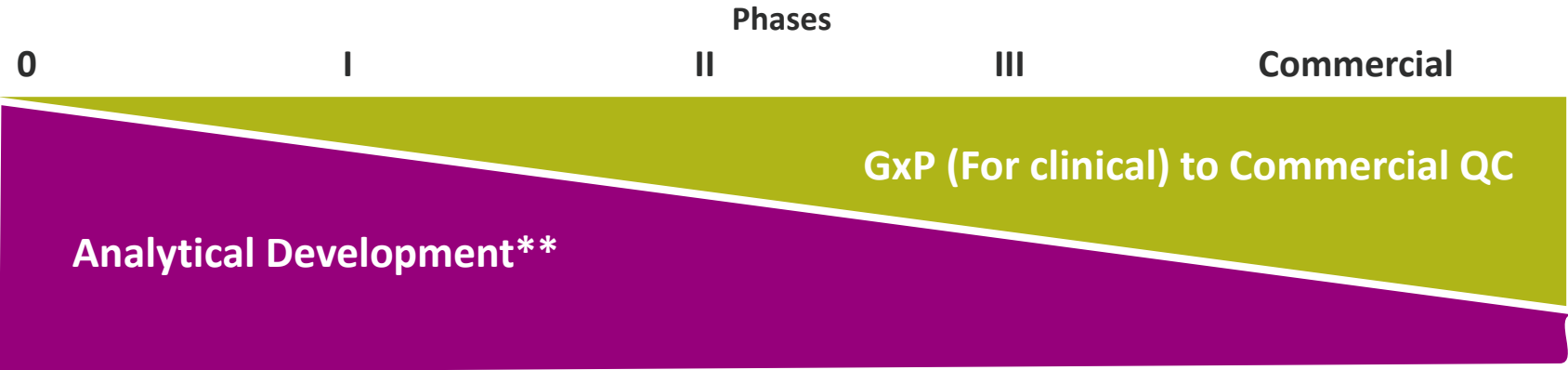
- :Develop scientifically sound methods
- :Methods transfer
- ::Methods characterization
- :Process Dev Support and R&D In-process testing
- :Product & process characterization
- : Determining Critical Quality Attributes
- :Development stability
- :Comparability studies
- ::Post approval improvements
- :Support regulatory filing

**Quality Control**

- :Good Mfg Practices (GMP) In-process testing
- :Release testing
- :Stability studies
- :Methods validation
- :Testing investigations
- :Method Life Cycle
- :Process Validation support
- :GMP studies
- : Support regulatory filing

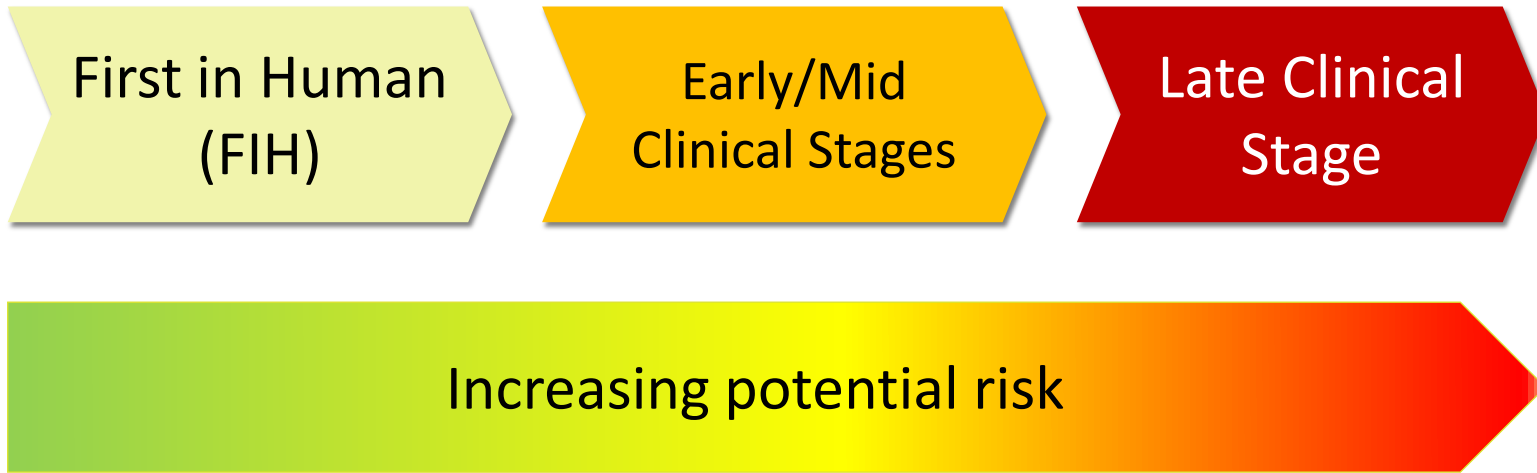
Close Collaboration is key:

Tech Dev  
 Analyt. Dev.  
 Manuf,  
 QC, QA  
 CMC Reg  
 CDMOs, CROs



\*\*AD work may be sustained post approval for PMC, Process improvements or trouble shooting purposes  
 PAV = Phase Appropriate Validation; PA = Phase Appropriate; with rapid CMC consideration

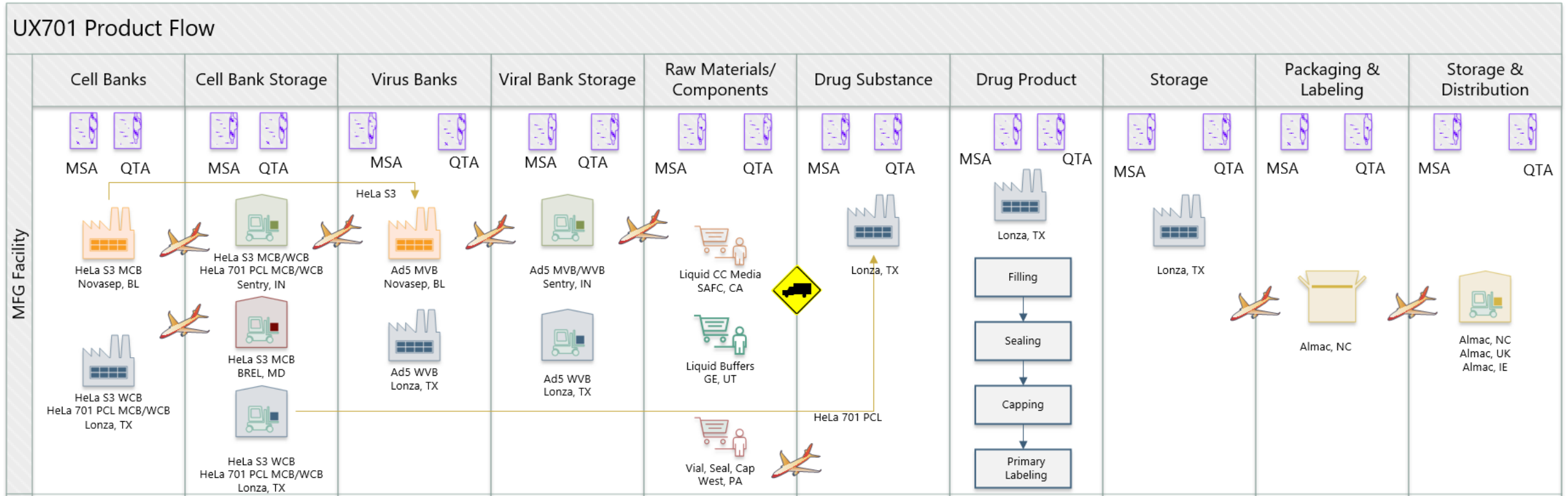
# CMC Risk Associated with Clinical Stage of Development



## ICH Q5E: Product comparability by Clinical Stage

Prior to Clinic	Not required
Early Clinical Stage	Not extensive
Mid Clinical Stage	More comprehensive
Late Clinical Stage	Comprehensive

# Supply Chain Can Be a Critical Success Factor



External Manufacturing requires a lot of management attention and oversight





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Questions?



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# Thank You



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