



Reneljix

Pharma Solutions

December 2023

Small Molecule APIs

R&D & Analytical Suite

Finished Dosage Facilities (USA)

Packaging Sources

Distribution & Market Access Centers

Key Highlights

- Manufacturing, process development, scale-up and optimization of novel APIs
- Solid State Characterization and API assessment
- Alternative Partners available in USA

- GLP labs
- R&D support for NMEs (New Molecular Entities) and Analytical Method Development.

- 5+ Billion Doses Manufactured/year
- 200+ Finished Dosage Formulations
- Ability to manufacture across all scales (Phase I-IV and Commercial)

- Dedicated site for manufacturing and supply of bottling, caps, and jars
- Dedicated site for pharmaceutical packaging
- Specialized marketing and label design services

- Specialized containment suites
- Network of 20+ depots globally
- Complete team of 200+ to provide support for market access

Facilities



Hyderabad, Telangana, India

500,000 sq feet



Hyderabad, Telangana, India

15,000 sq feet



Hauppauge, NY, USA

180,000 sq feet



Central Islip, NY

300,000 sq feet



Central Islip, NY

Windsor, NJ



NJ, NY, TN

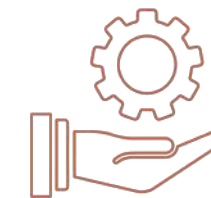
350,000 sq feet

20+ Clinical Depots Located Globally

CDMO Services

CPO Services

Support Services



Service offering goods

Development

Formulation & Drug Development

Sourcing

Scale up

Tech Transfer

Process Analytics Development

API production

Extraction

Synthesis

Fermentation:
Small Molecules

Other methods

Formulation

Solids

Semi-solid

Non-sterile liquids

Packaging

Primary packaging
(e.g., bottle)

Secondary packaging
(e.g., box, carton)

Tertiary packaging
(e.g., barrel, container)

Marketing & Label
Design Services

Included

Distribution 20+ Depots

Market Access and
Pricing Agreement with
Government Agencies

Regulatory Filing &
Support (IND, ANDA,
NDA, MAA)

Portfolio and Post-Life
Cycle Management

Scale

Small-scale production (preclinical - phase IV)

Large-scale production (phase III, phase IV, commercial)

- ✓ Early-Stage Drug Development
- ✓ Clinical Supply (Preclinical- Phase III)
- ✓ Commercial Manufacturing
- ✓ Dispensing for patients
- ✓ Generics/OTC
- ✓ Oral Solid Dose
- ✓ Oral Solutions & Suspensions
- ✓ Softgels
- ✓ Topicals
- ✓ Contract Manufacturing
- ✓ Packaging/Labeling
- ✓ Regulatory Support & Services
- ✓ PK/PD Modeling
- ✓ Taste Masking
- ✓ Bioavailability Enhancement Technologies
- ✓ Solubility Enhancement
- ✓ Customized Release
- ✓ Pediatric Formulations
- ✓ Poor Acceptability
- ✓ Dysphagic proof design
- ✓ Dosing Flexibilities
- ✓ Chemical Incompatibilities
- ✓ Orally Disintegrating
- ✓ Dosing Convenience
- ✓ Controlled Substances
- ✓ Abuse Deterrence Testing
- ✓ Oral Disintegration
- ✓ Customized Release
- ✓ Controlled Release
- ✓ Patient Centric Dosage Design



Precision Oral
Delivery



Minitablets



Taste Masking
Solutions



Softgels



Capsules



Orally
Disintegrating
Tablets



Modified Release
Forms



Tablets



Multi-
particulates



Oral
Suspensions



Visit renejix.com/oral-technologies to learn more

Key benefits of the integrated Early to Late offer



Enhanced speed

Proactive assessment and transfer allows for paralleled work



Knowledge Transfer

Avoided rework due to aligned systems and data sharing



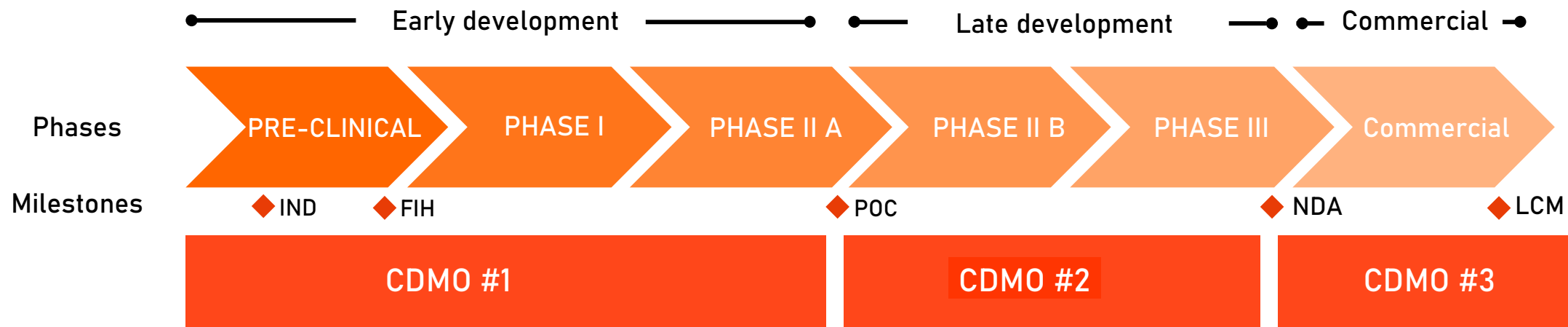
Reduced Staff Time

Through working with one partner point of contact that coordinates across phases



Cost Savings ~47%

Stay with one CDMO partner that has capabilities to support development from early to late stages



Multiple CDMOs


Cycles through CDMOs that specialize in different development stages

Meet our Experienced Team

Team of experts with **20-30+ years experience** in oral dosage development and commercialization

Integrated Business Development and Organizational Structure focused on quality and time to market

Strategic Group Leadership



Madhava Reddy CEO & President
Global Pharma

Business Leadership & Operational Team




Vineeth Reddy
Director – Business Development
Hauppauge, NY
Central Islip, NY



Ravindra Konakanchi
Head of Operations
Hauppauge, NY




Ragotham Reddy
Senior Director – Operations & Manufacturing
Central Islip, NY




Sridhar Gumadeveli VP – Research & Development
Hauppauge, NY
Central Islip, NY




Arti Mahajan
Director – Procurement & Supply Chain
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Venkat Jayaraman
Head of Quality Assurance
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Sunil Khan
VP – Head of Quality & Regulatory
Central Islip, NY
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Raj Vidap
Head of Analytical Sciences & Quality Control
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
Mohit Reddy
Program Manager
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Central Islip, NY



Rajesh Patel
Production Manager
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Narender Uppugalla
Sr. Director – Formulation
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Phani Jaggavarapu
Head of Analytical Sciences & Quality Control
Central Islip, NY