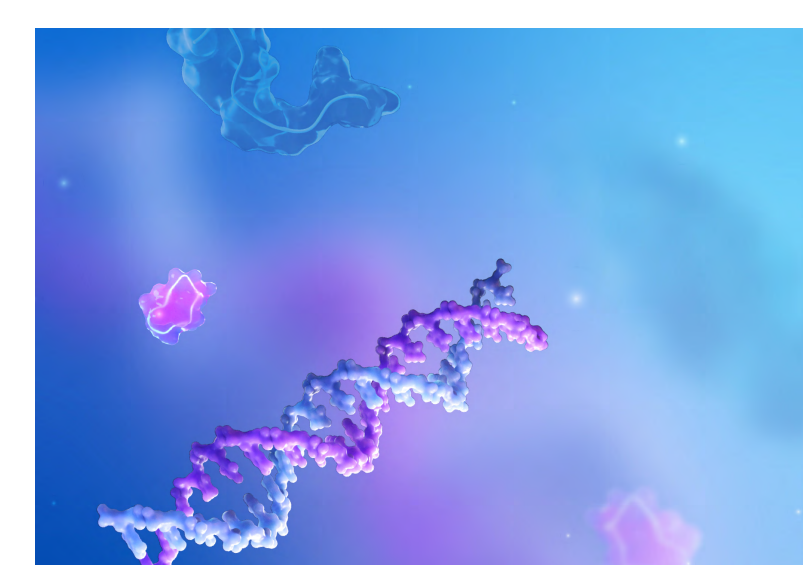


End-to-End CRDMO Platform

for Oligonucleot de and Peptede Therapeutics

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About Us

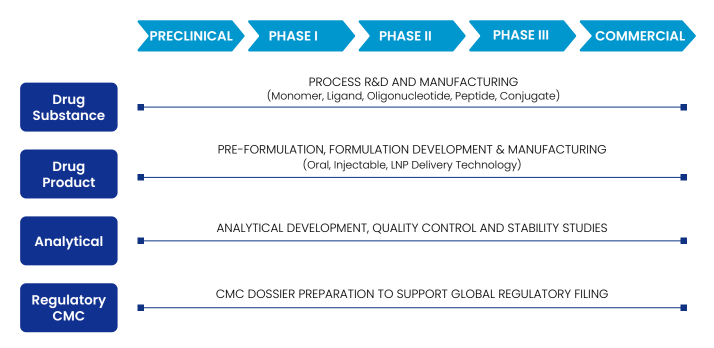
WuXi TIDES, a leading Contract Research and Development Manufacturing Organization (CRDMO) platform, is an integral part of WuXi AppTec. WuXi TIDES offers our worldwide partners efficient, flexible, and high-quality solutions for the drug development of oligonucleotides, peptides and related synthetic conjugates ("TIDES" drugs). We greatly simplify the TIDES drug development by providing all discovery, CMC development and the entire manufacturing supply chain under one roof. With over 1,100 scientists from 10 R&D and manufacturing sites, we offer discovery compound screening and synthesis, process development and manufacturing of novel monomers, linkers and ligands, oligonucleotides, peptides and complex synthetic conjugates at any scale. Beyond chemistry, we offer formulation development, manufacturing, labeling and distribution services in a variety of injectable dosage forms and filling formats including the Lipid Nanoparticle (LNP) drug delivery platform. Our comprehensive analytical method development, validation and testing platform will support your needs in TIDES drug development from discovery through clinical to commercial. Moreover, our Regulatory Affairs CMC team is experienced in preparing CMC dossiers to support global filings for TIDES new drug applications.

Trusted Partner

End-to-End CRDMO Platform for Oligonucleotide and Peptide Therapeutics

Oligonucleotide Chemistry Platform from Discovery to Commerical	Monomer/Ligand • Amidite • mRNA NTP • GalNAc • Linker, Spacer, CPG	Oligonucleotide • DNA • ASO • PMO • siRNA • sgRNA • Aptamer • microRNA • gRNA • Degenerate Oligonucleotide Pools	Conjugate • GalNAc- oligonucleotide • Lipid-oligonucleotide • 'Drug'-oligonucleotide • Dye-labelled oligonucleotide
Peptide Chemistry Platform from Discovery to Commerical	Unnatural Amino Acid • α,β,γ-substituted AAs • N-alkylated AAs • Glyco AAs, etc.	Peptide • Long Linear Peptide • Cyclic Peptide • Modified Peptide • PEGylated Peptide • Peptidomimetics	Conjugate • Peptide- Oligonucleotide (POC • Peptide-PMO (PPMO) • Peptide-Drug (PDC) • Radionuclide-Drug (RDC)

Oligonucleotide and Peptide CMC Platform



WuXi TIDES R&D and Manufacturing Network

Discovery Site



Shanghai, China discovery oligonucleotide & peptide, preformulation and formulation development

API Development and Manufacturing Site



Changzhou, Jiangsu, China API process development and manufacturing

Tianjin, China discovery oligonucleotide, amidite, GalNAc



Chengdu, Sichuan, China discovery peptide, unnatural amino acid



Wuhan, Hubei, China discovery peptide



Taixing, Jiangsu, China API process development and manufacturing



Singapore API process development and manufacturing (Operational in 2026)

Formulation Development and Manufacturing Site



Wuxi City, Jiangsu, China formulation development and manufacturing



Middletown, DE, USA formulation development and manufacturing



Couvet, Neuchâtel, Switzerland

Drug product manufacturing

OligonucleotideP Peptide



▶ 03 WuXi TIDES

Maintaining the Highest Global Standards

WuXi TIDES has an ingrained quality culture and follows the same quality and EHS system across all sites globally, with proven track record of inspections from all major regulatory agencies and our global customers.



14 US FDA 2013 - 2023



6 EMA 2009 - 2023



50 China NMPA 2015 - 2024



9 Japan PMDA 2019 - 2024



4 South Korea MFDS



4 SwissMedic 2018 - 2023

320+ Client audits every year

Why WuXi TIDES

Scalability

From discovery to development and commercialization all within WuXi TIDES with readily available large R&D and manufacturing capacity

Conjugation Chemistry

Seamless collaboration among oligonucleotide, peptide and small molecule teams

New Technology

Oligonucleotide: Biocatalysis for gRNA synthesis, Thin Film Evaporation (TFE) Peptide: Reactor-in-series (with PAT data collection), continues flow chromatography, Tangential Flow Filtration (TFF)/precipitation

Global Quality Standard

One quality system across all sites approved by major regulatory agencies around the world

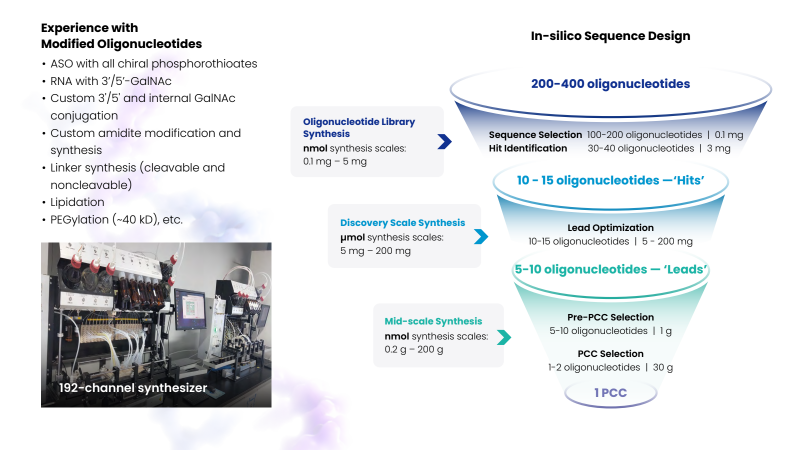
Comprehensive Analytical Platform

Method development and validation, IPC and release testing, characterization, stability

Integrated CMC

API process R&D and manufacturing, formulation development and manufacturing, analytical, CMC dossier preparation and clinical supply services

Oligonucleotide Discovery Services



0.1 mg-200+ g Synthesis Capability Produce 50,000+ Oligonucleotides Per Year

Up to 158-nt Long Oligonucleotides Access to 300+ Modifications & Conjugations

Comprehensive

Extensive experience in various modalities

Diversity

Large variety of linkers/conjugation strategies

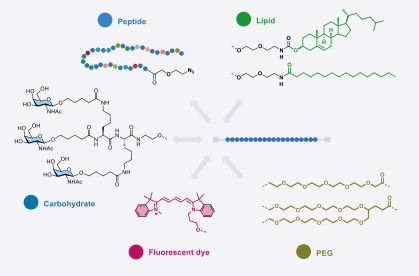
Flexibility

Custom-tailored solution according to your application

Linker Chemistry

Topology

Amide coupling Thiol-maleimide addition Click chemistry: CuAAC | SPAAC Disulfide chemistry Hydrazone chemistry **3'- or 5'-end conjugation Oligo moiety conjugate on** Base Phosphorus backbone **Complex dual conjugation:** 3',5'- | 3',3'- | 5',5'-



Peptide Discovery Services

High-Throughput Peptide Discovery Synthesis

mg to kg scale High quality – Over 98% Success rate High purity – Up to 99.9% Purity Fast delivery – 2 weeks turnaround time for most peptides under 30 AA at mg scale

Customized Peptide Synthesis

Up to 200 AA modified peptide with ligation Up to 70 AA modified peptide with SPPS only Modification: Dye/Biotin | PEG | Isotope labelling Peptidomimetics | Peptoid | PNA Cyclic peptides: Thioether | Disulfide | Biocyclic | Lactam | RCM | Lactone | Click

Our quick turnaround time and greater than 98% success rate ensures that our customers advance their projects quickly and efficiently. Most peptides under 40 amino acids are completed within 2 weeks from order placement with MS and HPLC/UPLC analytics data. Flexible deliverables include on-resin, crude, as well as purified powder at desired purity up to 99%.



Automatic Synthesis and Purification Platform

Automated solid-phase (Fmoc) synthesis provides efficient and reliable custom sequences up to 160 AA with over 80% purity. Our platform is equipped with a wide range of instruments to fit project requirements with a greater than 95% success rate.

Broad range of instruments

- CEM liberty blue
- Biotage Syro II
- CSBio automatic synthesizer
- Gilson Prep-HPLC
- Agilent UPLC
- Symphony X

Peptide Drug Conjugate (PDC) Platform

With our Integrated HPAPI capabilities, large linker library, and comprehensive chemistry platforms, we support Peptide Drug Conjugate (PDC) development from discovery to commercial.

Served 350+ customers and

delivered 30,000+ peptides in 2023

2,500+

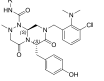
Unnatural amino acid catalog products

Bicyclic peptide

RCM monocyclic peptide

(Double RCM ring also achievable)

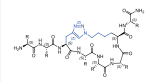




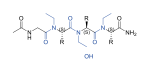
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PNA Length up to 30

Click chemistry



Peptoid



Oligonucleotide API Process Development and Manufacturing

We have 27 lines that cover a variety of Oligonucleotide types including ASO, siRNA, Aptamer, Oligonucleotide conjugates, PMO and PPMO.

- Utilize various modified monomers (2'-H, 2'-OH, 2'-F, 2'-OMe, 2'-OMOE, LNA, cEt, LNA, Vinyl-phosphate, Spacer, etc.)
- Chiral oligo synthesis
- Modification at 3'-end or 5'-end with GalNAc, triphosphate, cholesterol, saccharides, peptides, maleimide, etc.

Our team supports oligonucleotide production from lab scale to commercial scale with more than 20 small- to midscale production lines and 4 large scale production lines up to 6.0 mol per synthesis run.

Morpholino Oligonucleotide (PMO) Development and Manufacturing

We have developed high loading solid-phase PMO synthesis process that can achieve >50% yield, >75% crude purity, >90% final product purity with <0.045 EU/mg endotoxin.

Our chemistry knowhow enables various PMO 5' modifications with improved cleavage & de-protection process. Our optimized process offers freedom to operate without IP concerns.

We have customized PMO synthesis reactors up to 100 L to support large-scale production. Our team has successfully completed kilogram scale PMO development and manufacturing projects.

Recent Experience

- Completed 100 batches of 900 mmol DNA production (>400 kg delivery in total) including PPQ enabling studies and PPQ campaign in 9 months
- Completed 3 x1.6mol ASO production in 2 months
- Completed 64 siRNA projects with modifications at 5'-end or 3'-end such as GalNAc and cholesterol modifications, including 30+ integrated API/DP projects



Peptide API Process Development and Manufacturing

Same as oligonucleotide, our peptide platform is also in Changzhou and Taixing Site with industry-leading capabilities and capacities, covering a wide range of peptides and their conjugates.

- Long linear peptides
- Cyclic/Bicyclic peptides
- Modified peptides
- PEGylated peptides
- Peptidomimetics
- Branched peptides

We have built extensive experience with peptide-based conjugates including PPMO, Oligo-Peptide, Peptide-Toxin, Peptide-Antibody etc.

We currently have 32,490 L total reactor volume of the Solid-Phase Peptide Synthesis (SPPS), with reactors up to 3,000 L. More capacities will be added in Taixing site and Singapore site.



Our dedicated oligonucleotide and peptide analytical team has more than 180+ scientists, supported by more than 400 co-located analytical scientists from the process analytical and QC team to ensure the capacity required.

A full suite of state-of-the-art analytical instruments, including two-dimensional UPLC coupled with high-resolution MS for sequence and impurity analysis, enables the safety assurance and reproducibility required for regulatory submissions.





General Required Tests for Oligonucleotide and Peptide

- Purity and assay
- Microbiology safety including endotoxin and bioburden
- Stability study

Oligonucleotide

Unique Capabilities

- Identity by MW and sequencing with 2D UPLC coupled Q-TOF, MALDI-TOF and TOF HRMS
- Backbone composition identification by 31P NMP
- Purity by QTOF HRMS and quadrupole LC-MS

Peptide

- Amino acid analysis and enantiomeric purity
- Identity by peptide mapping
- Identity by MW and sequencing with 2D UPLC coupled Q-TOF, MALDI-TOF and TOF HRMS

Comprehensive Analytical Testing

Formulation Development and Manufacturing Platform

Drug Product Overview

- Pre-formulation
- Formulation development
- Clinical & commercial manufacturing

Sterile Parenteral Formulation Manufacturing Platform

- Disposable bags
- Mixing-filtration-filing with sterile
- Containers in a fully isolated system
- Wholly automated, robotic operation
- High potency (HP) injectable drug manufacturing (Can handle OEB5 compound; OEL: 10 ng/m³)



Dosage Forms

- ISolution/Emulsion
- Suspension
- Lyophilized powder
- Liposome/LNP
- Hydrogel

Filling Capacity

• Vial, prefilled syringe, cartridge

Lipid Nanoparticle (LNP) Platform

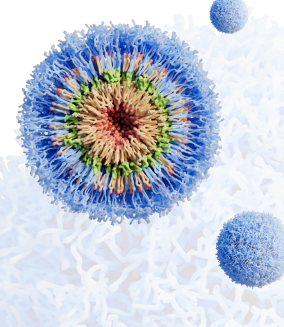
- Novel multi-channel mixing n-port provides robust scalability and reproducibility
- Novel lipid design and synthesis at any scale

Research & Development

- Robust scalability and reproducibility (start from 5 mL)
- Small particle size (80~100 nm) and narrow PDI (< 0.10)
- Optimized ultrafiltration and sterile
 filtration process

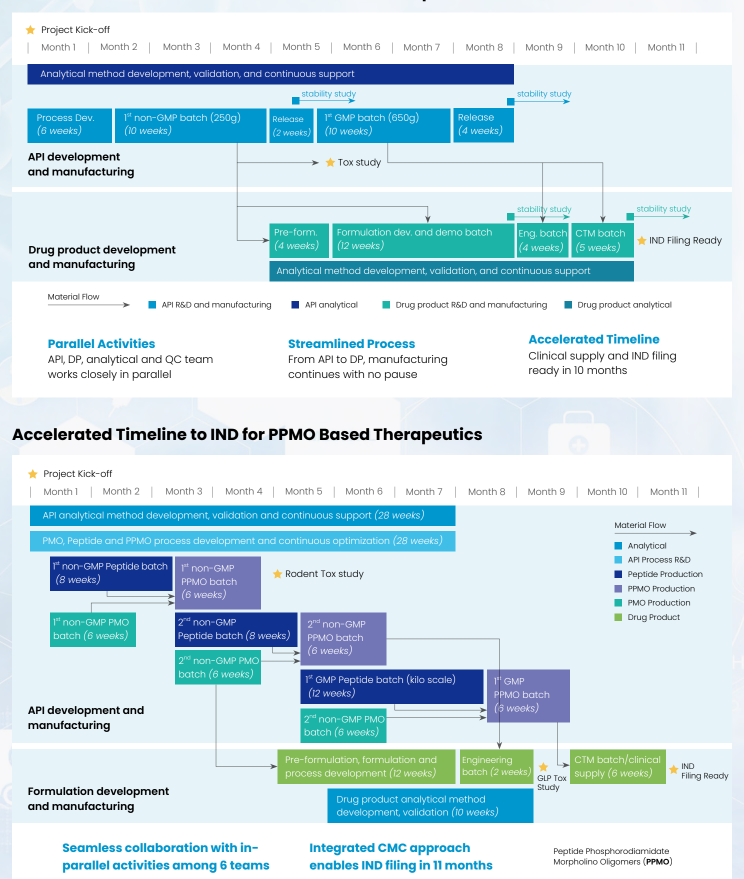
GMP Manufacturing

- Modular designed
- Flexible scale, 0.5 L 50 L per sub-batch

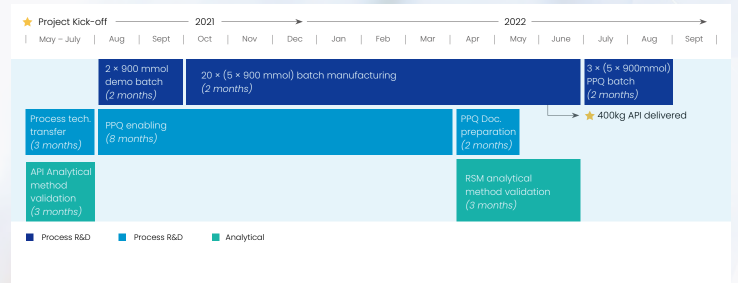


Case Study

Accelerated Timeline to IND for siRNA Based Therapeutics



Tech. Transfer and Commercial Manufacturing of Single Strand DNA



Quality and Speed

Successfully completed tech transfer, PPQ enabling, PPQ campaign, and commercial production (>400kg delivery) in one year

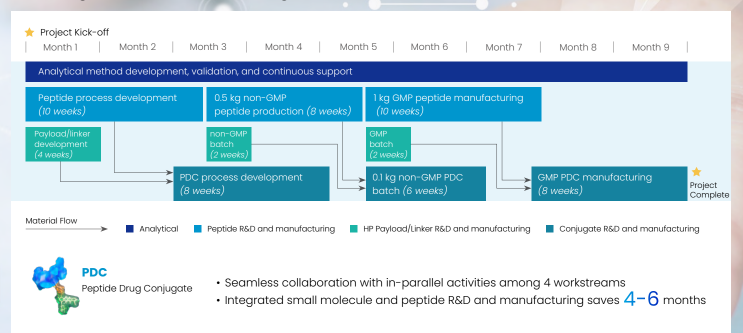
Continuous Improvement

Reduced commercial production cycle time by 50%

Robust Supply Chain

Established robust supply chain covering all raw materials, reagents, solvents, etc. used in oligonucleotide commercial production

PDC Drug R&D and Manufacturing in 9 Months



Regulatory Affairs CMC Platform



CMC submission packages written to support global IND and NDA filings from 2019 to 2023

Streamlined CMC writing integrated as part of the project team



Project Initiation

Provide RA consultation for phase and modality appropriate, country specific project scope and strategy

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Project Execution

Provide filing template, collect data once testing completed. Finish section writing along different phases and perform timely data review with clients to mitigate potential risks



Project Completion

Complete submission-ready CMC dossiers

Phase appropriate RA CMC filing strategies tailored to oligonucleotide and peptide drug characteristics

Adopt principles of ICH Q3 (impurities) and Q6 (specifications) to oligonucleotide and peptide based on our experience with O&P IND and NDA



10+ Global ESG Recognitions





Dow Jones Sustainability Indices Powered by the S&P Global CSA









- · Conducting formal process safety evaluations prior to manufacture
- Providing thorough health and safety training programs for both employees and contractors
- · Conducting toxicological and risk assessments for all new introduced processes
- Working with our suppliers and customers to minimize environmental impacts across the entire production and supply value chain
- Reducing our overall usage of water, energy, waste production, and emissions

ESG

REGIONAL

· To enhance our delivery of EHS policies, we have created several bespoke groups to drive forward a culture of corporate citizenship, including; EHS committee; process safety management committee; general manager representative; and a standalone EHS department



Our promises to health, safety and environmental care

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