





THE POWER TO MAKE







WHAT DO YOU WANT TO MAKE?





Ajinomoto Bio-Pharma Services gives you The Power To Make

To make your vision a reality. To make your program a success.

To make a positive difference in the world.

As a fully integrated CDMO, we provide therapeutic drug process development and manufacturing services to biotechnology and pharmaceutical companies worldwide.

With locations in Belgium, the United States, Japan, and India, we leverage our global infrastructure to provide the adaptive solutions, responsive service, support, and guidance needed to achieve your goals.

What truly differentiates us is our dedication to quality and our commitment to fostering trusted partnerships.

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Together, we have the Power To Make.



Your programs deserve the most comprehensive suite of CDMO services available, and Ajinomoto Bio-Pharma Services has the Power To Make your therapeutic vision a reality — from pre-clinical through commercial production.



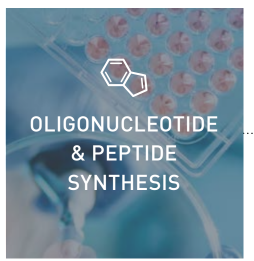
- Small Molecules
 - Lab & Pilot Scale Facilities
 - Dedicated HPAPI Labs
 - PAT/QbD Employed
- Large Molecules
 - Microbial Cell Strain Selection
 - Fermentation
 - Recovery & Purification
 - Complex Formulation Process Development
 - Terminal Sterilization Cycle Development
- High Potency/ADC Conjugation & Purification
- Oligonucleotides & Peptides



- High Potency API Units
- Cryogenic & High Pressure Reactions
- High Energy Chemistries
- Biocatalysis
- Continuous Flow Technology
- Chromatography
- Controlled Substances (EU)
- Life Cycle Management of APIs



- Protein and Plasmid Production
- Microbial Cell Banking & Characterization
- Upstream: Fermentation / Product Recovery
- Downstream: Purification / Formulation
- Corynex® Protein Expression System
- High Potency/ADC Purification & Conjugation



- AJIPHASE® Synthesis Technology
 - DNA/RNA Oligonucleotides
 - Phosphorodiamidate Morpholino Oligonucleotides (PMO)
 - Long Chain Peptides
- Small Scale Solid Phase Synthesis
 - Custom Oligonucleotides
 - Long RNA Chains
 - Long Chimera DNA/RNA Chains



- Aseptic Formulation
- Fill & Finish Capabilities
 - Aseptic Vial Filling
 - Aseptic Syringe Filling
 - Lyophilization
 - Terminal Sterilization
- High Potency/ADC: Fill & Finish and Lyophilization
- Packaging, Serialization & Aggregation Services



- On-Site Laboratories for In-Process & Lot Release Testing
- Phase Appropriate Analytical Method Development & Validation
 - Product Characterization
 - Reference Standard Qualification
 - Comparability Studies
- ICH-Compliant Stability Programs



PROJECT MANAGEMENT SUPPORT

At Ajinomoto Bio-Pharma Services, our project management team is focused on meeting the needs of our clients and their products. From RFPs through product shipping and all the milestones in between, the project management team works to balance the needs and expectations of all stakeholders while focusing on the end goal: high quality products that help patients.

DEDICATION TO QUALITY

We are committed to providing our clients quality products and services from pre-clinical to commercial programs into late life cycle, and any phase in between.

At every site, our experienced quality management team provides robust QC, QA, and compliance systems using shared best practices and a routine review of operational metrics to actively drive process optimization.

With our strong focus on quality, we offer our clients peace of mind and knowledge that their products will be a continued success. We take pride in knowing that our dedication is helping countless patients for years to come.

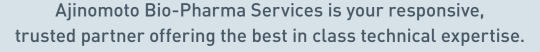
GLOBAL REGULATORY EXCELLENCE

Ajinomoto Bio-Pharma Services maintains an impressive and successful regulatory track record. Our quality management team focuses on continuous improvement, ensuring we are inspection-ready at all times — whether it's a regulatory agency or a customer inspection.

Our regulatory support team has contributed to numerous clinical and marketing application submissions, in addition to Drug Master Files, and is ready to help with your regulatory submissions, clinical filings, and commercial launches.



A GLOBAL DIFFERENCE



As a member of the Ajinomoto Group, we offer an extensive global reach with facilities in Belgium, United States, Japan, and India.



Over 40 years CDMO experience



1200+ employees worldwide



Excellent quality and regulatory history



Bio-Pharma manufacturing operations in 4 countries

WITH AJINOMOTO BIO-PHARMA SERVICES, YOU HAVE THE POWER TO MAKE

You have the power to make a difference by delivering new therapeutics that improve quality of life and inspire a healthier world.

You need a manufacturing partner who has the power to make your every challenge their own and who shares your unwavering tenacity and dedication from pre-clinical programs through to commercial successes.

Together, we have the Power To Make.

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