Your Biologics CDMO

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About Us

With a history spanning 17 years, Mabion has a wealth of experience in developing and manufacturing of biologic drugs, allowing us to meet the needs and requirements of the most demanding clients. Along with extensive bioanalytical capabilities and expertise in sterile manufacturing, packaging and serialization, we offer complete, end-to-end CDMO services.

Our Quality Management System, covering GMP, GLP, GCP and ISO, has been inspected by multiple authorities, assuring that services delivered by Mabion satisfy all regulatory requirements.

FACILITIES

Konstantynów Łódzki Facility

Mariana Langiewicza 60 Str., 95-050 Konstantynów Łódzki, Poland

GMP, ISO-certified

Manufacturing
Development
Analytics

Clinical, Commercial Process, Analytical methods Analytical/QC services for GMP/non-GMP product testing, incl. Cell Based Assays

Quality Regulatory

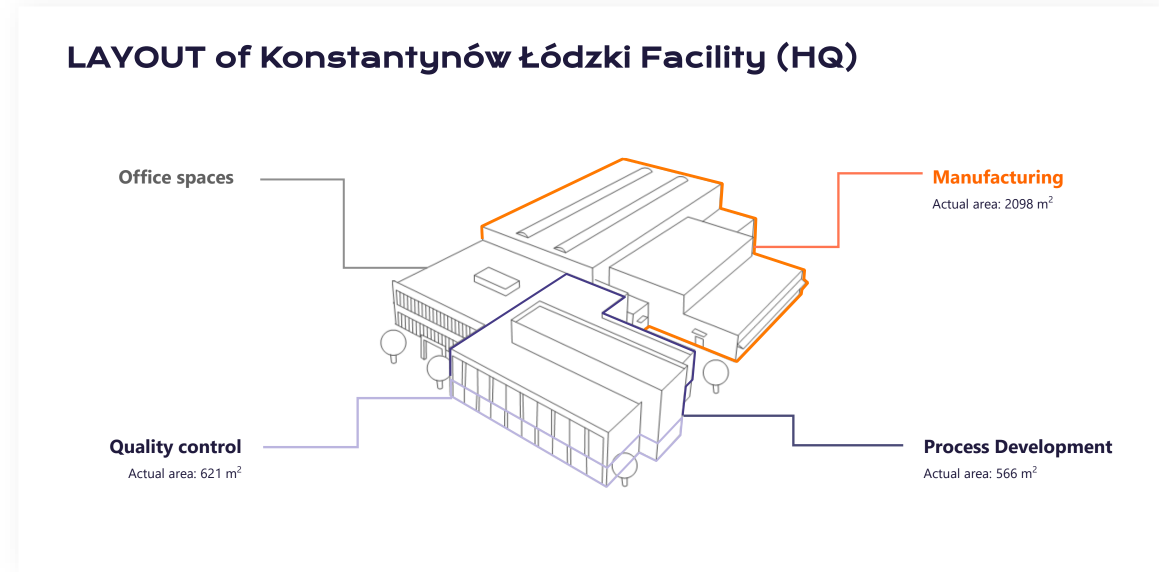
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Łódź Facility Fabryczna 17 Str., 90-344 Łódź, Poland

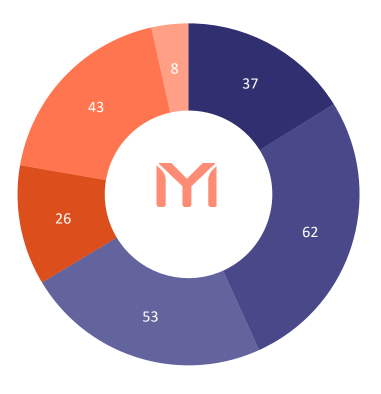
GLP-certified

Bioanalytical studies Clinical trials PK, PD, Immunogenicity; BSL-II labs Design, Operational support

Warsaw 1,5 h from airport to HQ



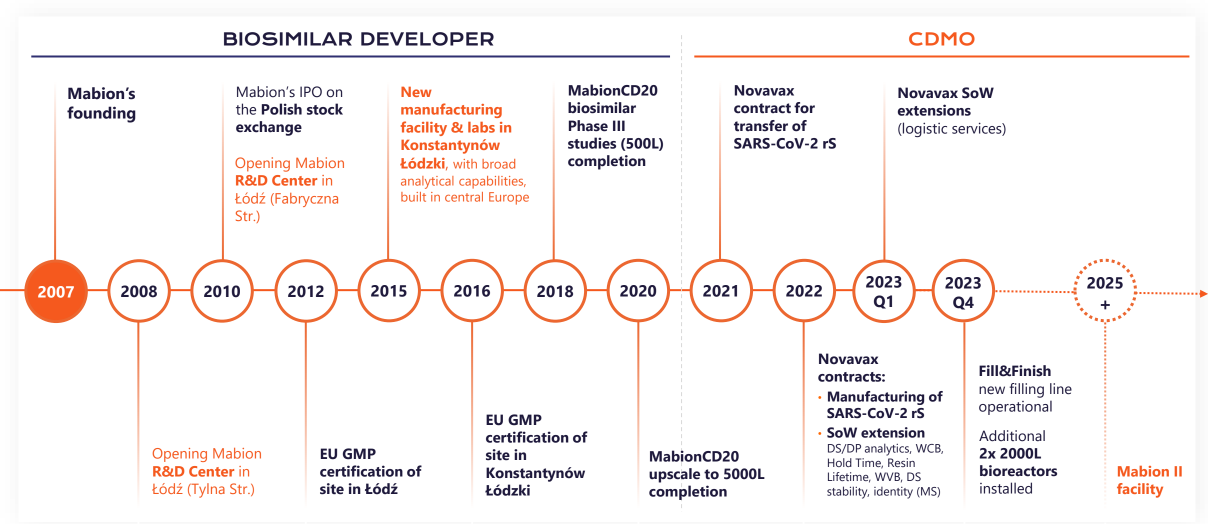
Team - 229 FTE's





History & Transformation into CDMO

History and future of Mabion



By developing own biosimilar products in the past, Mabion has acquired key competencies and assets, building an integrated biopharmaceutical company

Thanks to these key competencies and assets, Mabion seized a market opportunity and since 2021 has been transforming into a CDMO



We have developed advanced competencies in biologic drug technology using cell lines and monoclonal antibody engineering for development, manufacture and control of biosimilars

We have developed effective processes that allow us to systematically obtain products of **high quality** within agreed timelines

We have achieved a high level of integration and we offer a **broad spectrum of services** in the areas of protein development, analytics and manufacturing, as well as consulting and regulatory advisory services

We have a **dynamic team with strong interdisciplinary experience**, competence to operate under GLP/GMP and an open approach ('can do' attitude)

We have modern analytical and manufacturing assets located in the EU (Poland)



We operate in compliance with the highest quality standards in the industry: GMP, GCP, GLP, ISO

building competence and resources

We have validated our competencies and we have begun to monetise the resources we have built through our first commercial collaboration

transformation into a biologics CDMO

2021



2024

2007

Biologics CDMO

Quality

Quality systems operating at Mabion include EU-GMP for manufacturing (since 2012), GLP for bioanalytical studies (since 2012) and ISO. Mabion QMS was built following EudraLex vol. 4 principles.

As a result, robust GMP processes have been established, ready to accommodate any Client's quality requirements, including compliance with the US FDA cGMP

Mabion's QMS

Current Mabion facilities regulatory status – GMP, GLP, GCP and ISO compliance		
Konstantynów Łódzki	Good Manufacturing Practice (GMP)	 Good Manufacturing Practice (GMP) is a system for ensuring that products are consistently manufactured and controlled according to quality standards It is designed to minimize the risks involved in pharmaceutical production that cannot be eliminated through testing the final product GMP certificate was granted in April 2017 for Konstantynów Łódzki facility - Scientific and Industrial Complex of Medical Biotechnology (Previously, in November 2012, for the Research and Development Centre in Łódź)
	ISO ISO	 Mabion holds three ISO certificates: 14001:2015 environmental, ISO 45001:2018 work safety regulations, ISO 50001:2018 energy management Audits were performed by independent certified specialist SGS Polska / SGS UK / SGS Italy Certificates were issued in 2023 for 3 years period
Łódź	Good Laboratory Practice (GLP)	 GLP defines a set of rules and criteria for quality system management of research laboratories in order to ensure the trustworthiness of laboratory data, including bioanalytical data from clinical studies and preclinical studies during drug development Mabion was granted GLP certificate in March 2014 and has been continuously re-certified every 2 years (recent GLP certificate is from 2024) Holding this certificate indicates that studies and analyses carried out at Mabion meet high international quality standards
	Good Clinical Practice (GCP)	 GCP defines the rules that constitute the international quality standard for clinical trials involving humans Compliance with GCP standards guarantees credibility and authenticity of the data collected during clinical trials All trials conducted by Mabion to date have been in accordance with GCP

Mabion audits and inspections history

- Since its founding in 2007, Mabion has passed multiple inspections and audits demonstrating compliance with GMP, GLP and GCP practices as well as ISO 9001 and ISO 14001/45001/50001 standards.
- > Quality assurance is subject to rigorous and continuous improvement through internal and external audits.
- > Mabion became GMP and GLP compliant in 2012. No critical findings were ever identified.

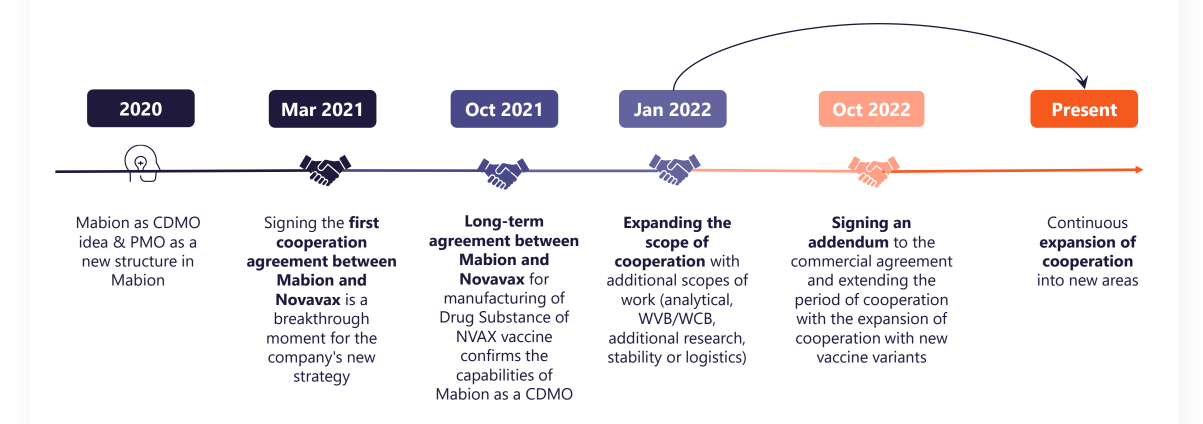


* Including 2 inspections performed at sites participating in a clinical study sponsored by Mabion.

Biologics CDMO

Novavax case study

A path from an idea to becoming a company providing commercial services in diverse biotechnology fields



Notable accomplishments in the Novavax lab-scale, analytics and process transfer

Mabion successfully finalized the feasibility phase during which the Novavax protein production process was transferred and scale-up to our facility **within 3 months and ahead of schedule**.

During this period, Mabion has accomplished:





Successful lab-scale batches



Successful full-scale batches Transfer of DS release testing analytical methods



Generation of >100 documents (SOPs, summary reports, etc.)



The entire process, from agreement signing to the final report and client approval, **took 30 weeks**

Notable accomplishments in the Novavax manufacturing of drug substance for COVID-19 vaccine

Novavax - Mabion Commercial production is a success with further extension of the scope of cooperation as well as future business development activities. Mabion has been able to adjust the work and schedule for Novavax's needs in short term and jointly solve process and analytical challenges.

Batch success rate and manufacturing schedule adherence per value stream and production suite were assessed on **100% in the KPI Analysis** performed by Novavax.

Until now, Mabion has accomplished:





100% successful engineering and transfer batches

Successfull completion of PPQ batches



GMP production of DS of SARS-CoV-2 in 2,500L scale started No failed batches and safety events



KPI scorecard review showed no safety events, great batch success rate and schedule adherence

Mabion - A trusted CDMO partner

The best testimony to our quality and reliability as CDMO is the recommendation issued by Novavax based on a 3-year history of successful collaboration on the protein COVID-19 vaccine.

"

"Mabion demonstrated flexibility and a high level of customer focus at the time when the Omicron variant arrived, as they managed to swiftly adapt the manufacturing process to the production of a modified vaccine antigen. This seamless transfer of technology and prompt commencement of the production for a new variant highlighted Mabion's agility and technical prowess."

"

"

"(...) based on the outstanding results of our cooperation, we can enthusiastically recommend Mabion as a trusted and reliable CDMO for the development and manufacturing of vaccines. The exceptional capabilities, state-of-the-art technologies and commitment to quality make Mabion an invaluable partner for any company wishing to outsource their key process.

"

"

"Mabion is fully capable of delivering this wide panel of services, while continuing to demonstrate a high level of professionalism and unwavering commitment to quality.



MABION Biologics CDMO

Your End-to-End Biologics CDMO Partner

Drug Substance Manufacturing Gene To Vial: End-to-End Clinical Service

Analytics

Fill & Finish

Process Development Cell Line Development & Banking



Services

Drug Substance Manufacturing

Fill & Finish

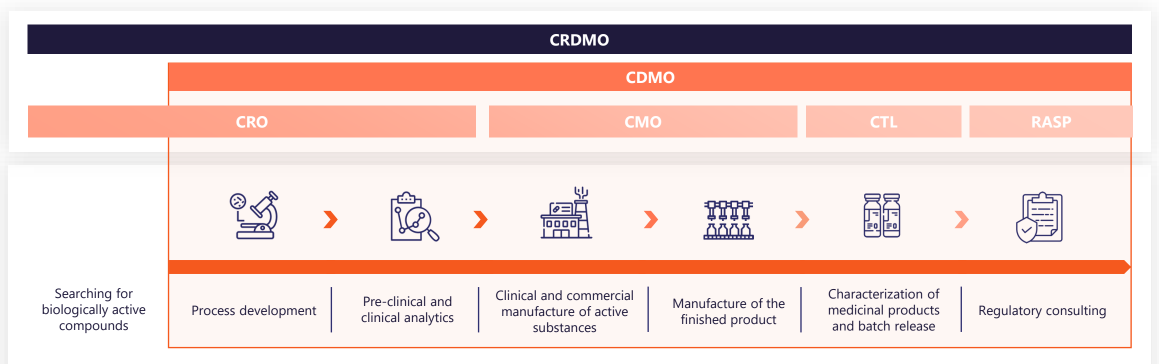
Process Development Gene to vial: End-to-end Clinical Services

Analytics

Cell Line Development & Banking

Mabion offers a comprehensive portfolio of services for a wide range of biological products

As an integrated CDMO, Mabion offers a full range of services, with a focus on recombinant protein technologies and antibody format, within which Mabion has all needed assets and is ready to execute commercial orders



> Mabion has an offering that addresses a wide range of the CDMO service value chain

> High level of integration and ability to address only selected stages from the entire CDMO service value chain (e.g., CMO, CTL, RASP, part of CRO)

CTL - Contract Testing Laboratory CMO - Contract Manufacturing Organization CDMO - Contract Development & Manufacturing Organization RASP - Regulatory Affairs Service Provider CRO - Contract Research Organization CRDMO - Contract Research, Development & Manufacturing Organization

Project Management

With every project entrusted to Mabion, comes a dedicated project manager. This committed person ensures that your project is given the utmost attention.

Our approach to **project management** is the key component of Mabion's commitment to provide a world-class, customeroriented outsourcing experience. By fulfilling this commitment, we are capable of delivering the top-quality services at competitive prices.

Fill & Finish

Gene to Vial: End-to-End Clinical Service

Process Development

Analytics

Cell Line Development & Banking

Drug Substance Manufacturing

> Mammalian & insect cell cultures

- > 2 x 2000 L, 2 x 200 L, 2 x 50 L stirred-tank, single-use bioreactors from Cytiva
- > 2 x 2500 L and 4 x 250 L orbital shaking bioreactors
- Medium & supplements preparation and storage capacity

 Separation technologies (depth filtration & centrifugation)

- > Affinity chromatographies lon-exchange chromatographies
- D O W N S T R E A M P R O C E S S

UPSTREAM

PROCESS

- Ultra/diafiltration
- Nanofiltration
- > Sterile filtration
- > Formulation
- > Buffer preparation



Fill & Finish

Gene to Vial: End-to-End Clinical Service

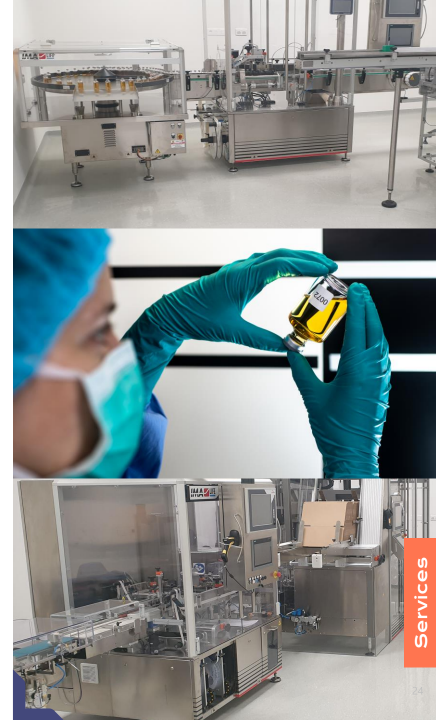
Process Development

Analytics

Cell Line Development & Banking

Fill & Finish

- > Automated filling line
- > Automated product inspection
- > Secondary packaging
- > Product storage and transportation
- > Serialization



Fill & Finish

Gene to Vial: End-to-End Clinical Service

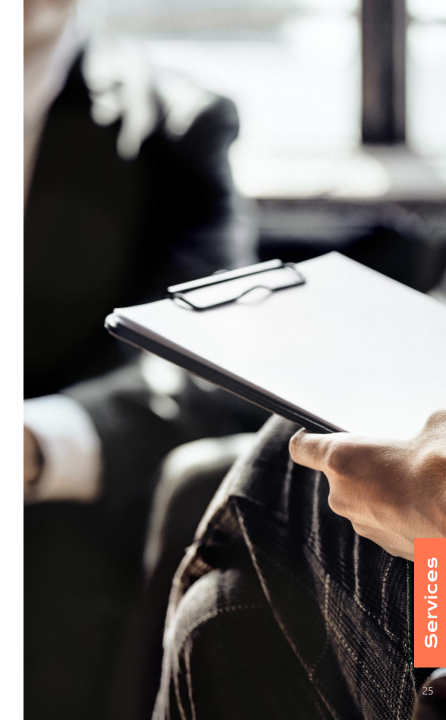
Process Development

Analytics

Cell Line Development & Banking

Gene to Vial: End-to-End Clinical Service

- > Cell Banking
- > Process development
- > Analytical development
- > DS manufacturing
- > Fill & Finish
- > Suite of services tailored to the project



Fill & Finish

Gene to Vial: End-to-End Clinical Service

Process Development

Analytics

Cell Line Development & Banking

Process Development

- > Upstream process development
- > Downstream process development
- > Process space & process characteristics
- > Process scale up
- > Analytical methods development & validation
 - > Structural assays
 - > Physicochemical assays
 - > Biological/functional assays
- > Comparability & similarity assessment
- > Reference standard establishment
- > Clinical and pre-clinical analytics development





Fill & Finish

Gene to Vial: End-to-End Clinical Service

Process Development

Analytics

Cell Line Development & Banking

Analytics

- Drug Characterization Services >
 - **Physiochemical Analytics** >
 - **Structural Analytics** >
 - **Biological Analytics** >
 - **Clinical and Preclinical Analytics** >
- GMP release testing >
- QC testing of intermediate product, drug substance, drug Product, > reference product
- Analytical methods development and validation >
- Comparability and Similarity studies
- Characteristics of reference standard >
- Long term, accelerated and stress stability study >
- Environmental monitoring >



Fill & Finish

Gene to Vial: End-to-End Clinical Service

Process Development

Analytics

Cell Line Development & Banking

Cell Line Development & Banking

- > Cell Line Development
 - > Clone Selection
 - > Stable and highly productive monoclonal cell line
 - > Culture medium optimisation
- > Non-GMP Research Cell Banks
- > cGMP Master Cell Banks
- > cGMP Working Cell Banks



Thank you

MABION

SCIENTIFIC AND INDUSTRIAL COMPLEX FOR MEDICAL BIOTECHNOLOGY

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Click here to watch a video presenting Mabion:



Link: https://www.youtube.com/watch?v=2hzQI5ZGyxk&t=2s