



Precision in Every Dose,  
Innovation in Every Form







## Pioneers in precision and innovation, dedicated to delivering excellence in every formulation.

BPI Labs is a FDA registered drug manufacturer and 503B outsourcing facility, providing the absolute highest quality medications to fill unmet needs. BPI services hospitals, clinics, and pharmacies throughout the US. As a leading drug manufacturer and 503B, we ensure the highest quality standards in every product we produce.

BPI Labs steadfast commitment to quality, is shaping the future of healthcare, one dose at a time.



## About Us

BPI Labs, LLC, a leading manufacturer of branded and generic injectable medications is proud to be one of a handful of FDA registered drug manufacturer and 503B. BPI's strong quality culture, led by experienced professionals adhering to the strictest GMP regulations, is the cornerstone of our operations. Using cutting-edge isolation and human less robotic manufacturing technologies and all in-house analytical and microbiological testing, we maintain the absolute highest quality sterile injectable production to meet market demands.

- FDA Registered
- 22 Approved Drugs
- Over 100,000 sq ft Sterile Manufacturing
- Over 200 Employees





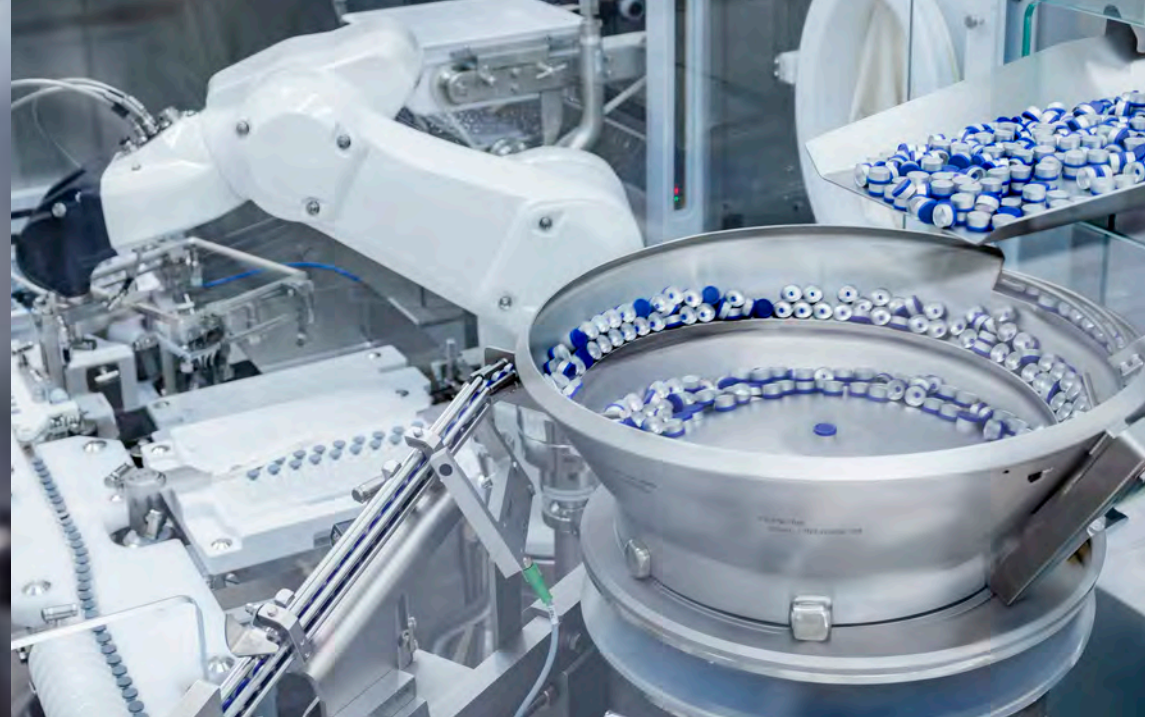
# Our Facilities

BPI's is one of the most technologically advanced sterile manufacturing facilities in the country. The facility was designed with cGMP adherence in mind.

- Fully Human-Less Robotic Filling Technology
- Automated Visual Inspection
- Sterile Water for Injection Manufacturing Plant
- 500 Liter Batch Capacity
- One-Time Use, Disposable Pathways
- In-House Analytical Labs
- In-House Microbiological Labs
- In-House R&D
- In-House Validation; IQ, OQ, PQ













## Quality

BPI's 503B drugs are made under the strictest cGMP guidelines, uses the same commercial manufacturing equipment and follows the same quality systems and procedural standards as its other FDA approved drugs. BPI's quality is far beyond reproach and is unequivocally the most reliable compounded drugs on the market. Our most recent FDA audit of our 503B resulted in approvals with no 483 observations.

## Exceptional Testing

Every batch is fully tested in our full chemistry and microbiology labs within the facility to meet more strict, self generated guidelines and parameters. This provides full confidence in producing industry best results for our consumers through strict compliance with cGMP regulations. We do not rely on third-party manufacturers' C of A to assure quality. All ingredients are tested in-house, upon arrival and must meet our strict standards and methods developed with our in-house QC chemistry lab and microbiology labs.

## Quality

BPI Labs only uses DMF active ingredients along with validated inactive ingredients to assure the purest materials are incorporated.

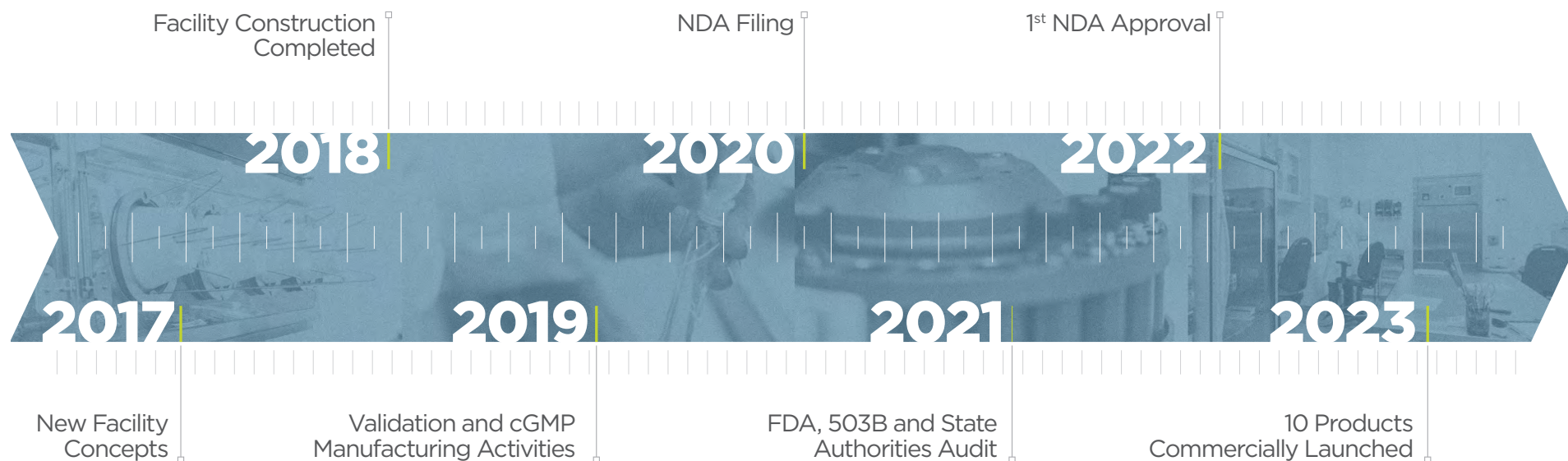
## Exceptional Testing

BPI Labs proudly adopts robotic technology and system into sterile pharmaceutical manufacturing and testing process, making its products unmatched in the generic injectable industry.





# Our Journey



## Capacities

Batch sizes range from 40,000 - 100,000 units and current capacity is in excess of 150,000 vials a week. Additional expansion is underway, which will increase capacities in excess of 2 million vials per month.

**22**  
Approved Drugs

**100,000+**  
sq. Sterile Manufacturing

**200+**  
Employees





## Our Difference

	503A (Compound Pharmacy)	503B (Outsourcing Facility)	BPI Labs (503B + FDA Drug Manufacturer)
<b>Regulatory Agency</b>	State (State Board of Pharmacy)	Federal (FDA) and State (SBOP)	Federal (FDA) and State (SBOP)
<b>Regulations</b>	USP <795>, <797>, <800>, SBOP	FDA 21 CFR Part 210 and 211 (cGMP)	FDA 21 CFR Part 210 and 211 (cGMP)
<b>Distribution</b>	Patient Specific Only	May or may not be. Can be sold to pharmacies or Office Use	May or may not be. Can be sold to pharmacies or Office Use
<b>Batch Size</b>	250 units	Limited; usually up to 25 liters	Unlimited; up to 500 liters
<b>Raw Material</b>	Need C of A	Require C of A and Manufacture with FDA registration	Only use DMF Material, validate, qualify and create STMs
<b>Quality Systems</b>	Pharmacist review their own work	Quality department in place	Full quality assurance department, independent of operations
<b>Batch Testing</b>	No Analytical Testing	Minimal; Outsourced	Full in process testing; All in house
<b>Stability Dating / Beyond Use Date (BUD)</b>	Literature Only	Minimal; Outsourced	Full stability protocol to establish BUD and expiration dates
<b>Validation (Methods)</b>	Literature	Outsourced; Minimal	Full validation
<b>Environmental Monitoring</b>	Every 6 Months	Per production batch	Exceed cGMP Requirements
<b>Endotoxin</b>	Required; Outsourced	Required; Outsourced	Required; In House
<b>Impurities Testing</b>	Not Required	Not Required	Required; In House
<b>Release Testing; Potency</b>	Not Required	Every batch; Outsourced	Every batch, including In Process; In House
<b>Visual Inspection</b>	Manual	Manual	Fully Automated; High Speed AI Equipment
<b>Sterile Filling</b>	Laminar Flow Hood	Laminar Flow Hood	ISO5 Robotic Aseptic Filling (Commercial)





## GLP-1 Products

With a steadfast commitment to excellence, we offer a product line that is meticulously crafted to uphold the highest scientific and quality standards. Our injectables provide customers with cutting-edge solutions to address the evolving demands of discerning patients. From innovative treatments to essential medications, each product undergoes rigorous testing and stringent quality control measures to ensure efficacy and safety. Partner with BPI Labs today and elevate your offerings with trusted injectables backed by unwavering dedication to patient well-being.

- FDA Registered Drug Manufacturer & 503B Outsourcing Facility License
- DMF (Drug Master File) Semaglutide Listed with FDA
- Ready to use Formulation
- Base Form Similar to Brand (No Acetate or Sodium)
- 360 Days Beyond Use Date (BUD)





# Semaglutide Injection

Product	Strength	Size	Container	NDC	Pack Size
Semaglutide Inj. 1 mg/mL	1 mg/mL	1 mL	Vial	54288-0826-25	25
Semaglutide Inj. 2.5 mg/mL	2.5 mg/mL	1 mL	Vial	54288-0833-25	25
Semaglutide Inj., 5 mg/2 mL (2.5 mg/mL)	5 mg/2 mL	2 mL	Vial	54288-0827-25	25
Semaglutide Inj., 12.5 mg/2.5 mL (2.5 mg/mL)	12.5 mg/2.5 mL	2.5 mL	Vial	54288-0831-25	25



# Tirzepatide Injection

Product	Strength	Size	Container	NDC	Pack Size
Tirzepatide Inj. 10 mg/mL	10 mg/mL	1 mL	Vial	54288-0835-25	25
Tirzepatide Inj. 30 mg/3 mL	10 mg/mL	3 mL	Vial	54288-0832-25	25
Tirzepatide Inj. 60 mg/3 mL	20 mg/mL	3 mL	Vial	54288-0836-25	25







BPI Labs helps to eliminate the logistical challenges of working with multiple compounding pharmacies, managing state licenses, and navigating capacity limitations, allowing you to focus on delivering exceptional care to your patients.

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