

FUJIFILM

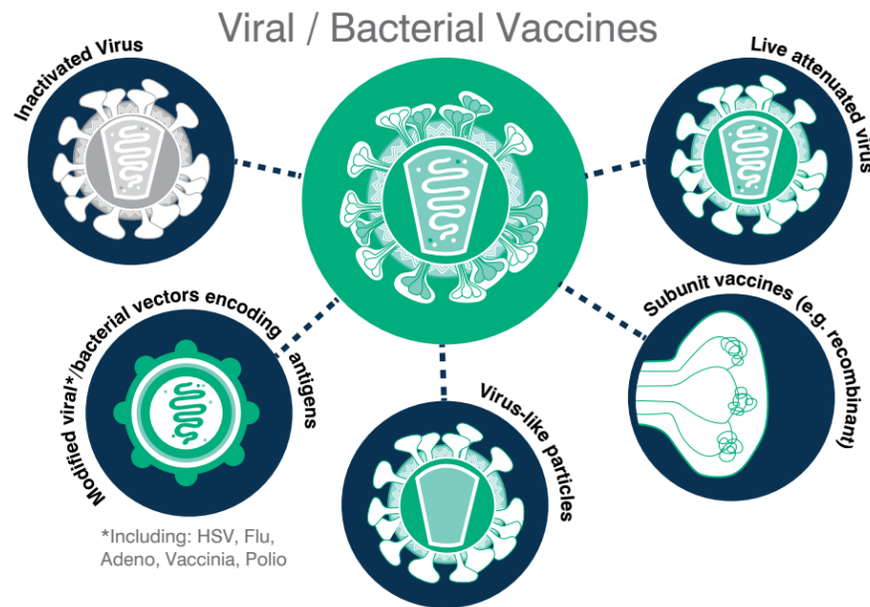
Diosynth
biotechnologies



VACCINE SERVICES

Your CDMO Partner **Advancing Vaccines** to Patients

Partners for *Life*
Advancing tomorrow's medicines



We are an established leader in advancing vaccines from preclinical development to commercial supply across multiple modalities.

End-to-end Vaccines Development and Commercialization

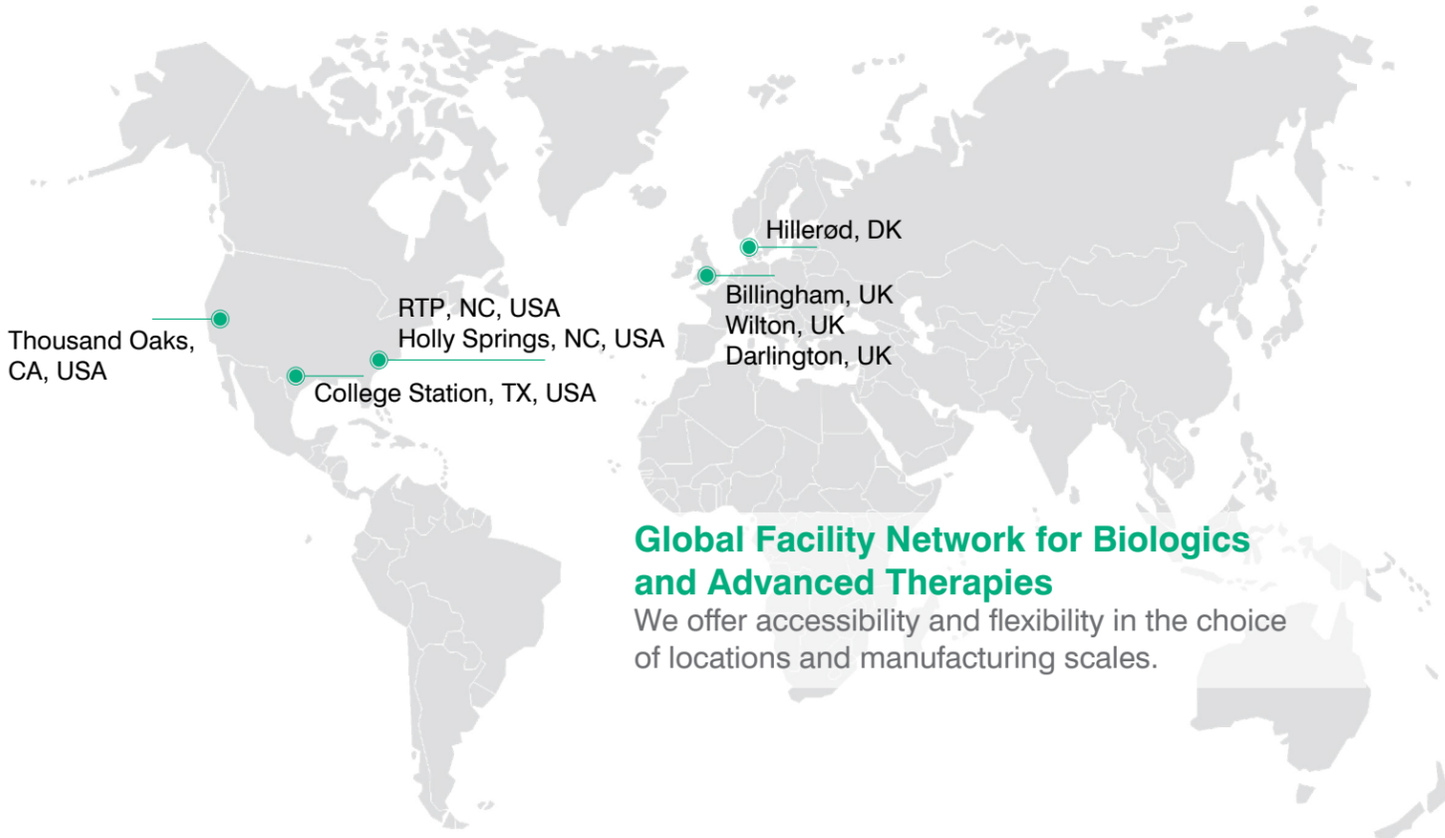
	Preclinical	Clinical	Commercial
cGMP cell banking and storage	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Process Development	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Toxicology material generation non GMP and GMP	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>
Large scale cGMP manufacturing	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
Product quality testing	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
Stability testing	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
Sterility and endotoxin	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
Process Characterization	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
Process Validation	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>

Vaccine Drug Product and Finished Goods Services

Development and manufacturing capabilities designed with patient safety and centrality in mind.

We use robotic systems with single use product path to meet your program needs, providing pre-clinical batches up to 10,000 units and commercial batches up to 150,000 units using RTU nested components in vial, syringe or cartridge format.

- Formulation development
- Stability testing
- Clinical and commercial DP manufacturing of vial, syringe and cartridge filling with state-of-the-art isolator technology
- Prefilled syringe (PFS) and vial labeling
- Primary packaging in single or multi-pack formats
- Secondary packaging and clinical labeling
- Assembly into auto-injectors and PFS components



[Contact us](#) to get the latest updates on our network and capabilities.



Contact us to discuss your science.

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