

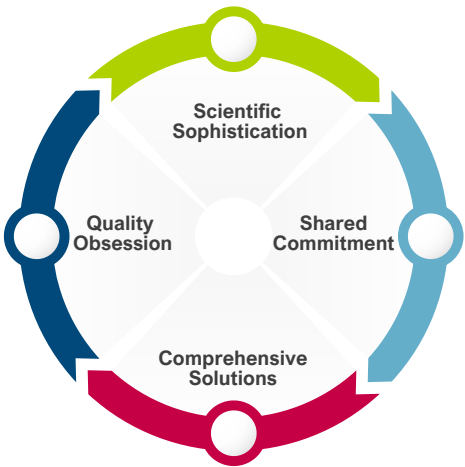


# Bring your innovation to its destination

with creative analytical solutions that advance your path to commercialization

## Who We Are

Solvias accelerates commercialization for innovative companies with CMC analytical solutions, leveraging deep scientific expertise and **a strong focus on customer success**. Our experts excel in **biologics, cell and gene therapies, and small molecules**, offering comprehensive services from raw materials to final product testing, including API development and manufacturing. With **six strategically located global facilities** and certifications in GMP, GLP, and ISO, we conduct over 100 successful audits annually, **ensuring science and quality drive faster, safer product development**.

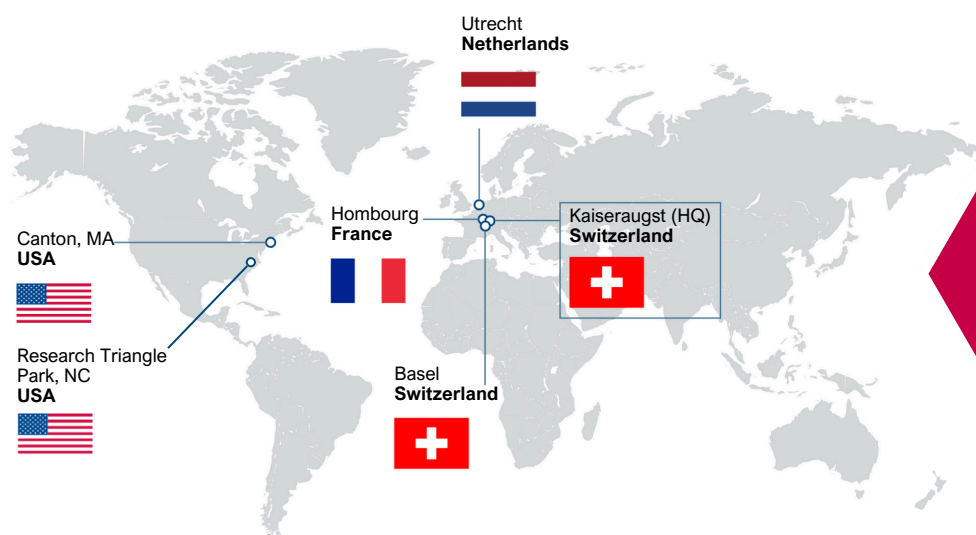


## Analytical Solutions that Span the Entire Product Lifecycle

	Research	Preclinical	Early-Phase Clinical	Late-Phase Clinical	Commercial
Analytical Characterization Services					
Cergentis Services - Genetic QC					
Small Molecule API Development & Manufacturing					
Custom Solutions					
cGMP Drug Substance & Drug Product Testing & Release					
Impurity Testing					
Biosafety Testing					
Excipient & Raw Materials Testing & Release					
Stability Studies					
Analytical Method Development					
Extractables & Leachables Testing					

## Why partner with us?

- CDMO/CRO
- Founded in 1999
- 800+ team members
- 175+ PhD-level scientists
- GMP, GLP, ISO9001 certified
- 22.5K sqm of lab capacity
- 700+ customers worldwide
- 6 centers of excellence



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“The Solvias team's expertise in genomic characterization has helped us establish the assays needed for each of our CRISPR genome-edited CAR-T cell therapies.”

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“The scientific delivery of this quality work from the Solvias team exceeded industry standards. The team are masters of their craft.”

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“The drug delivery and physical characterization team provided outstanding support to fine tune analytical methods, perform product specific characterization studies and to complete in vitro BE studies required for product filing at the US FDA.”

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**Contact us to speak with  
an expert:** [info@solvias.com](mailto:info@solvias.com)

