



ROADMAP to success

Provisetm Viral Clearance Services

Accessibility and availability of viral clearance services is critical when it comes to on-time filing with regulators. Finding a partner with the capacity to meet your needs in a timely fashion and superior quality can be a challenge.

The Evolution of Demand

Demand for Provisetm Clearance increased during the pandemic due to travel restrictions. From study design and tech transfer all the way through to support with regulatory questions, Provisetm Clearance offers:

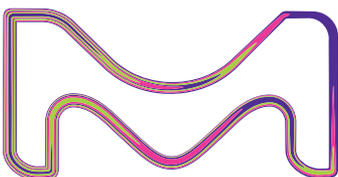
- Ability to focus on high value-added activities at your own facility
- Reduced risk: No need to stock virus in-house
- Cost savings including travel expenses

Our experience:

- Monoclonal antibodies
- Recombinant proteins
- Novel and biosimilar products
- Vaccines
- Gene therapy vectors

Team of expert scientists trained in:

- Virus reduction filtration
- Column chromatography
- Ethanol fractionation
- Viral inactivation by treatment with solvent/detergent and extreme pH
- Column re-use studies





Trust Provide™ Clearance Services

*We do it for you,
keeping you informed along the way*



Discuss your clearance needs
with an expert partner you can trust?
SigmaAldrich.com/provide

Understanding Your Molecule

Our Provide™ Clearance Services offer a team of expert process scientists with years of experience in viral clearance studies who work with you to design your IND- and BLA-enabling studies to assess your manufacturing's ability to remove or inactivate viruses.

Most importantly you will develop a close relationship with your dedicated team of experts, who will be in regular communication to keep you informed of your study's progress along the way.

Collaboration sessions include among others:

- Technical discussions to understand every detail of your process
- Advice on study design
- Preparing workbooks, SOWs and study designs
- Updates from your study including: Cytotoxicity, mock runs, and process steps

When the study is complete, you will receive a final report and support through your regulatory journey.

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At-a-glance

 **520**

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The cover image was produced before the Covid19 crisis started.
We take our responsibility seriously and fully comply with all protection rules.

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