

VIRAL CLEARANCE SERVICES

VALIDATION OF BIOPHARMACEUTICAL
PROCESSES FOR VIRUS ELIMINATION





The advent of the biopharmaceutical age has opened new avenues of disease treatment and prevention. However, biopharmaceutical products, such as monoclonal antibodies, recombinant proteins, vaccines, blood derivatives, and animal products carry an inherent risk of transmitting infectious viruses due to the source material used, manufacturing processes, and routes of administration. Several instances of viral contamination of biopharmaceutical products have occurred in the past. In order to control viral contamination of biological products, the following complementary programs are now set up by bioprocessors as a standard:



Selecting and testing the starting materials (cell banks, raw materials, etc.).



Testing of the product at various stages of production.



Incorporating viral inactivation, removal steps into the production processes, and testing the manufacturing process for viral removal or inactivation capacity (i.e., viral clearance studies).

Commonly used methods of detecting viruses include infectivity assay [50% tissue culture infectious dose (TCID50) and plaque assays], molecular probe methods [polymerase chain reaction (PCR), hybridization, western blot, etc.], electron microscopy, antibody production in animals, reverse transcriptase test, and immunoassay for viral-specific proteins, etc. However, direct testing alone is not sufficient to ensure viral safety due to the inherent limitations in all virus detection methods. For example, unknown viruses, certain virus variants, and viruses that were thought not to be potentially pathogenic can not be detected. In addition, low concentrations of virus may escape detection due to the sensitivity limit of the assays.

Because of these testing limitations, an evaluation of the production process for its ability to inactivate and/or remove a wide variety of known or unknown contaminating viruses is necessary to ensure viral safety of biopharmaceutical products.

OVERVIEW OF A VIRAL CLEARANCE STUDY

The goal of a viral clearance study is to assess the capability of the production process to remove and/or inactivate any potential viral contaminants. Experience and knowledge in selecting the most relevant target viruses and their surrogates, purification steps, and cleaning agents are critical in the design and implementation of a high-quality and cost-effective viral clearance study. It is also very important to “plan ahead” in order to ensure that the Sponsor’s timeline is met.

Generally, a viral clearance study, from initial planning to final report issuance, is comprised of the following steps:

- ◆ Risk assessment of the product regarding viral safety
- ◆ Selection of target viruses to be tested for viral clearance
- ◆ Selection of process steps to be evaluated for virus elimination
- ◆ Down-scaling the selected manufacturing steps
- ◆ Preliminary studies of cellular toxicity and viral interference
- ◆ Viral-spiking process runs
- ◆ Collection and titration of the process samples
- ◆ Data generation and calculation of viral clearance factors
- ◆ Issuance of the final report

RISK ASSESSMENT OF THE PRODUCT

Assessing the viral safety risks associated with a specific product requires the consideration of several factors, including type of starting materials, reagents used in the purification, manufacturing process, phase of clinical development, product indication, and specific patient population. This assessment is used to determine appropriate target testing viruses and the goals for the viral clearance study. As an example, for products derived from murine cell lines at the IND submission stage or for phase 1 clinical trial, it is usually sufficient to examine only the clearance of murine retrovirus. However, a comprehensive panel of at least four viruses would be necessary before phase 2 and 3 materials are made.



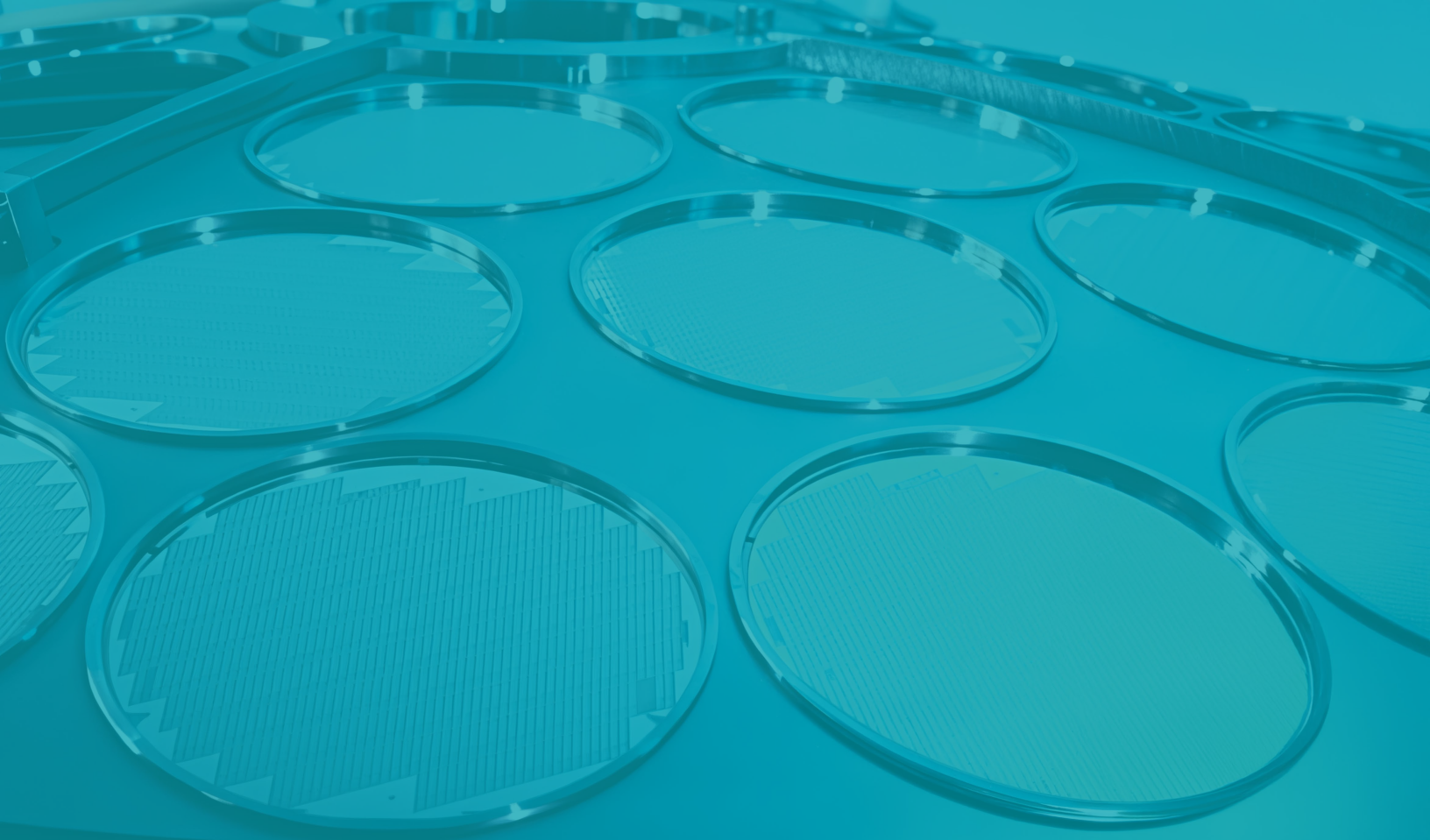
REGULATORY GUIDELINES FOR VIRAL VALIDATION STUDIES

Regulatory expectations on viral safety for biopharmaceutical products have evolved over the past several decades. Original concerns focused on a relatively small number of known viruses associated with the production cell lines. Today the concerns are much broader, encompassing unknown and uncharacterized agents. These increasingly stringent standards are intended to decrease the risk of transmitting viruses.

ICH Q5A, Viral Safety Evaluation of Biotechnology Products Derived from Cell Lines of Human or Animal 1 Origin , (1997) specifically requires that a manufacturer of biological products for human use demonstrate the capability of the manufacturing process to remove or inactivate known contaminants.

The FDA's Points to Consider in the Manufacture and Testing of Monoclonal Antibody Products for 2 Human Use (1997) and Points to Consider in the Characterization of Cell Lines Used to Produce Biologicals 3 (1993) apply the basic concepts of the ICH Q5A guideline and provide further description on viral safety evaluation methods. They provide a good general guideline for bioprocessors to consult when evaluating the viral safety of their products in development.

The 1996 CPMP guidelines on viral validation, Note for Guidance on Virus Validation Studies: The Design, Contribution and Interpretation of Studies Validating the Inactivation and 4 Removal of Viruses and CPMP's Note for Guidance on Plasma Derived 5 Medicinal Products , provide detailed recommendations for the manufacturers of biopharmaceutical products to follow when performing viral validations. These recommendations also set specific values for virus removal and inactivation levels that had to be attained.



With implementation of the EU Clinical Trials Directive 2001/20/EC 6, all EU Member States now require submission of an Investigational Medicinal Product Dossier (IMPD) starting at phase 1. The virus safety evaluation for biotech products is part of the IMPD's quality documentation.

The EU's requirements on viral clearance evaluation are further summarized in the EMEA Guideline on Virus Safety Evaluation of Biotechnological Investigational 7 Medicinal Products (2008), and two replicate evaluations per step are generally recommended.

Similar viral safety validation and testing strategies are described in the International Standard ISO 22442-3: Medical devices utilizing animal tissues and their derivatives – Part 3: Validation of the elimination and/or inactivation of viruses and 8 transmissible spongiform encephalopathy (TSE) agents (2007) for medical devices derived from animal material.

All of these regulatory guidelines emphasize that each viral validation study should be reviewed on a case-by-case basis and that log reduction factors obtained should be viewed under experimental limitations and product specific risk factors.



SELECTION OF VIRUSES FOR THE STUDY

Based on viral safety risk assessment of the product, a panel of relevant and model viruses should be chosen for the viral clearance study. Relevant viruses refer to the viruses that have been identified as contaminants or potential contaminants of the starting or intermediate materials. When the potential viral contaminant cannot be readily propagated or assayed in the laboratory, a model virus, i.e., a virus with properties similar to the agent in question can be used. In addition, the model viruses represent a broad spectrum of different physicochemical properties of viruses, so that the safety of the product from contamination by adventitious viruses is ensured.

Other biological and laboratory aspects of virus such as ease of achieving high titer, availability of a sensitive and robust assay system, and safety of staff when using high-titered virus should be considered in the selection of viruses for use in viral clearance studies.

Table 1 and Table 2 list the viruses commonly used in the clearance studies. In addition to the organisms listed, other viruses specifically related to the product may be chosen to meet the individual needs of the sponsor.

TABLE 1: VIRUS SELECTION FOR PRODUCTS DERIVED FROM MURINE HYBRIDOMAS AND CELL LINES

VIRUS	FAMILY	GENOME	ENVELOPED?	SIZE (MM)	RESISTANCE	MODEL FOR
XMuLV	Retroviridae	ss-RNA	YES	80-110	Low	Non-defective C type retrovirus
PRV	Herpesviridae	ds-DNA	YES	150-250	Low-Medium	Human or animal herpesviruses
PI-3	Paramyxoviridae	ss-RNA	YES	150-200	Low-Medium	Potential contaminant of bovine serum
MMV	Parvoviridae	ss-DNA	NO	~20	Very High	Human parvovirus B19
Reo-3	Reoviridae	ds-RNA	NO	60-80	High	Rotavirus, Bluetongue virus, orbiviruses

TABLE 2: VIRUS SELECTION FOR PRODUCTS DERIVED FROM BLOOD AND PLASMA

VIRUS	FAMILY	GENOME	ENVELOPED?	SIZE (MM)	RESISTANCE	MODEL FOR
HIV-1	Retroviridae	ss-RNA	YES	80-110	Low	HIV-1, HIV-2 and HTLV
PRV, HSV-1	Herpesviridae	ds-DNA	YES	150-250	Low-Medium	Human or animal herpesviruses
BVDV	Flaviviridae	ss-RNA	YES	40-60	Low-Medium	Hepatitis C Virus
BPV, PPV	Parvoviridae	ss-DNA	NO	~20	Very High	Human parvovirus B19
HAV, PV-1, EMCV	Picornaviridae	ds-RNA	NO	25-30	High	Hepatitis A Virus
HAV, PV-1	Caliciviridae	ss-RNA	NO	30-40	High	Potential contaminant of bovine serum

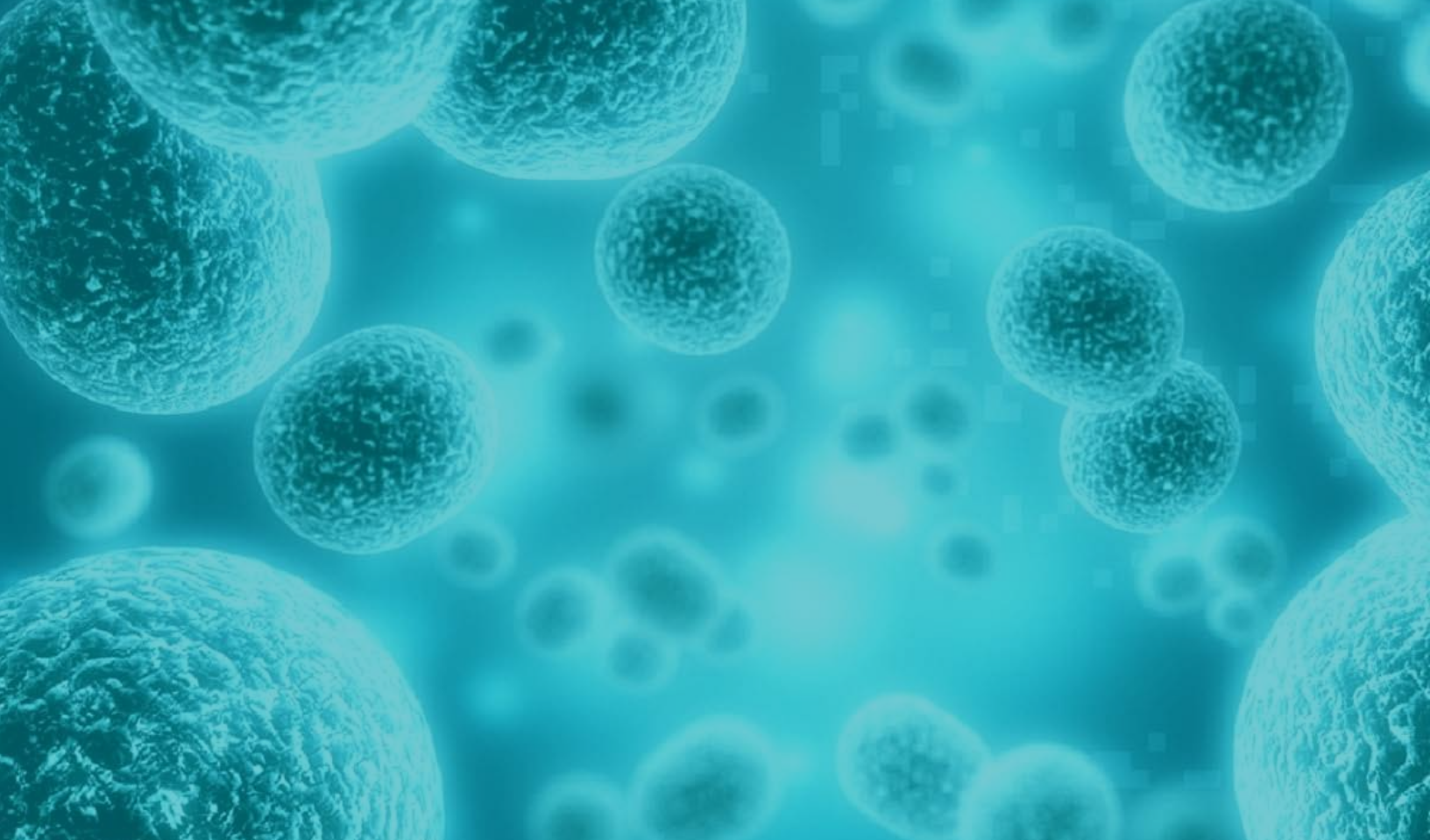
SELECTION OF PROCESS STEPS FOR EVALUATION

A satisfactory level of virus elimination for bioprocesses is usually achieved by a combination of inactivation of the virus and physical removal of the virus from the product (partitioning). The steps that can remove or inactivate viruses may be part of the actual purification procedure or have been specifically introduced for the sole purpose of viral clearance. Examples of different steps that can generate viral clearance are shown in Table 3.

TABLE 3: PROCESS STEPS COMMONLY EVALUATED FOR VIRUS REMOVAL AND INACTIVATION

VIRUS INACTIVATION		VIRUS REMOVAL	
Chemical Methods	pH	Chromatography	Affinity
	Detergent		Ion Exchange
	Solvent / Detergent		Hydrophobic Interaction
	Alcohol		Reverse Phase
	Disinfectants		
Physical Treatment	Heat	Virus Filtration	
	Pressure	Centrifugation	
	Irradiation	Precipitation	

A manufacturing process for biopharmaceuticals should incorporate at least two distinct robust virus clearance steps, with at least one step effective on non-enveloped viruses. Robust steps are those able to clear a wide range of viruses and are not influenced by process variables (pH, protein concentrations, buffers, temperatures, etc.). Other considerations for selection of process steps for viral clearance evaluation include avoiding the selection of steps with the same mechanism of clearance and ease of scaling-down of the steps.



DOWN-SCALING THE MANUFACTURING STEPS

Scaling-down the process steps to be evaluated is a prerequisite to performing the actual spiking experiments, as it would be impractical to use the actual production scale for the viral clearance study due to the volumes of virus needed. Also, it would be inappropriate to introduce high-level infectious virus into the cGMP manufacturing facility.

In order for studies performed on a laboratory scale to be extrapolated to the manufacturing scale, the validity of the scale-down must be proven. For an inactivation step, the pH, protein concentration, buffer, and temperature must be considered. For a chromatography step, the resin, bed height, linear flow rate, contact time, buffer, pH, ion strength, protein concentration, elution profile, and temperature must be considered. The overall level of purification (product purity and yield) is critical for any process scale-down.

In addition to process scale down, other issues such as column sanitization and reuse are important for a satisfactory viral clearance study. It needs to be demonstrated that when the chromatography columns are expected to be reused, the viral contaminants bound to the resin will be inactivated between production cycles and the viral elimination capacity will remain consistent throughout the life cycle of the columns.

PRELIMINARY STUDIES FOR CYTOTOXICITY AND VIRAL INTERFERENCE

In order to obtain accurate viral clearance data for a process step, the effect of the test samples (process intermediates) on the viral assaying system must be evaluated. Generally, clearance factor could be underestimated if production components are cytotoxic or viral interfering.

As most viral infectivity assays utilize mammalian cell lines as the host, the toxicity of the process materials on the host (indicator) cells is evaluated. Serial dilutions of the test sample are incubated with the indicator cells. Those dilutions that result in a change in cell morphology will be regarded as cytotoxic and excluded from the spiking experiment. The lowest dilution at which no cytotoxicity is observed is regarded as the non-cytotoxic dilution.

In addition to cytotoxicity, more subtle problems, such as samples interfering with the ability of the virus to establish infection in the host cells, may occur with the process samples. In many cases samples that show no signs of cytotoxicity can show significant viral interference. Serial dilutions of the test sample are mixed with a range of virus dilutions and inoculated onto the host cells. The infectivity titer obtained from test sample treated titrations is compared to the titer from cell medium-made titrations. The lowest dilution at which no interference is observed is regarded as the noninterfering dilution.

The cytotoxicity and viral interference studies are typically performed prior to the spiking experiments. The results are used to determine the sample dilution ratio in order to remove any cytotoxic and viral interfering effect of the collected process samples.



PERFORMING VIRAL-SPIKED PROCESS RUNS

After the preliminary studies are completed, the viral spiking experiment can be performed on down-scaled process steps. The starting material for each step is spiked with high-titered virus stock and processed. The spiked load, process intermediates, and product fraction (or post-treatment sample for inactivation studies) samples are collected, diluted with medium (to remove cytotoxicity and/or viral interference), and assayed. Worst-case conditions are used when process parameters are variable in order to illustrate the extremes of the viral clearance capacity.

For virus inactivation studies, the kinetics of inactivation needs to be examined by comparing the pre-, intermediate, and post-treatment samples. For virus removal studies, the distribution of virus should be studied by assaying the process intermediates in addition to the spiked load materials and the product fraction.



CALCULATION OF VIRAL CLEARANCE FACTORS

Upon collection and dilution (quench), the process samples are typically titrated immediately. If this is not possible and it is necessary to freeze samples before titration, a freeze/thaw virus control should be added. Serial dilutions of the process samples are inoculated onto host cells and incubated under appropriate conditions for a defined period of time. At the end of incubation, each well is scored for viral presence, typically by observation of viral-induced cytopathic effect (CPE). The virus titer of each sample is then calculated by methods such as the Spearman-Kärber method.

Viral titers are normally expressed with 95% confidence limits. The accuracy and reliability of the viral assaying methods are emphasized by all regulatory guidelines. Accuracy, reproducibility, repeatability, linearity, limit of detection, and limit of quantitation need to be demonstrated by assay validation.

The limit of detection of the infectivity assay is directly related to the volume of inoculum. Therefore, when virus concentration is low, assaying a larger volume of a test sample increases the sensitivity of the assay and in turn may increase the viral reduction factor able to be demonstrated. The large volume sampling is particularly useful for achieving a good reduction factor when the test samples have a high level of cytotoxicity, as the input viral load is limited by the titer of the challenge virus stock.

CALCULATION OF VIRAL CLEARANCE FACTORS

The viral load of the spiked starting material is compared to that from the post-processing material to calculate the viral reduction factor for the process step. The titer of each sample is multiplied by the volume of the sample and any post-collection volume manipulation (such as dilution or pH neutralization). The 95% confidence interval for the reduction is calculated based on the 95% confidence limits for the input and output samples. The log viral reduction factor for each process step can be added together to obtain the viral clearance value for the production process as a whole, however, it must be determined that the steps evaluated are independent (i.e., they utilize different mechanisms to eliminate the virus).

ISSUING FINAL REPORTS

After the viral clearance data are generated, Microbac Laboratories, Inc. will issue an unaudited preliminary results summary to the client within 2-3 business days. Then, the final report will be written and sent to the Quality Assurance (QA) unit for audit. The entire data package will accompany the final report.

When interpreting viral clearance results, emphasis is often given by regulatory groups to the robustness of the process steps rather than the actual clearance factor. CPMP guidelines require at least one robust step for recombinant murine products, and at least two robust steps for blood products^{3,4}. The ICH and FDA Points to Consider documents emphasize the robustness of the overall production procedure and require that for murine retrovirus, the level of clearance should be substantially in excess ($\geq 10^4$ - 10^6) of the potential viral load^{1,2}.

After completion of the QA review, the final report will be issued to the sponsor.



PROCESS VALIDATION AT MICROBAC LABORATORIES, INC.

Microbac Laboratories, Inc., is a contract antimicrobial testing laboratory serving biopharmaceutical companies in the United States and abroad. With a team of highly experienced staff members, worldwide experience, an internal database for viral clearance studies and our track record of exceptional client services and competitive pricing, we can help you design and execute high-quality and cost-effective viral clearance studies to meet your needs as well as regulatory requirements.

For further information, please call 703-925-0100 or email Sterling.CS@microbac.com.

VIRUS ABBREVIATIONS

BPV - Bovine Parvovirus

BVDV - Bovine Viral Diarrhea Virus

EMCV - Encephalomyocarditis Virus

FCV - Feline Calicivirus

HAV - Hepatitis A Virus

HIV-1- Human Immunodeficiency Virus Type 1

HSV-1- Herpes Simplex Virus Type 1

MMV - Murine Minute Virus

PI3 - Para-influenza Virus Type 3

PPV - Porcine Parvovirus

PRV - Pseudorabies Virus

PV-1 - Poliovirus Type 1

Reo-3 - Reovirus Type 3

XMuLV - Xenotropic Murine Leukemia Virus

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