Concentration Of The baculovirus Autographa californica M nucleopolyhedrovirus (AcMNPV) by ultrafiltration

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1 Abstract
Concentration and retention of a rod-shaped virus during tangential flow ultrafiltration (UF) was assessed to evaluate the potential of membrane-based downstream methods with advantages such as easy scale-up for industrial processes. A recombinant baculovirus of the non-spherical Autographa californica M nucleopolyhedrovirus (AcMNPV), vHSGFP, expressing egfp was filtered using polyethersulfone membranes ranging from 30 to 1000 kDa molecular weight cut-off (MWCO). A 20-fold virus concentration was achieved when a membrane cut-off range of 100 to 1000 kDa was tested. Fouling was observed and cake formation and pore plugging were postulated as concurrent causes with different impact depending on the MWCO. A reduction of virus concentration in the range of 2 to 5 log units in the permeate was observed illustrating the potential of membrane-based virus filtration as a useful unit operation in downstream processing.

2 Introduction
Virus clearance is critical in pharmaceutical and biotechnological processes [1]. Downstream processing of viruses as products (e.g., viral vectors expressing genes for gene therapy) requires virus concentration and purification [2]. However, virus purification using sucrose or cesium chloride density
centrifugation is time consuming and results in significant virus particle losses [2]. Pelleted viruses from centrifugation can be highly viscous and difficult to resuspend in solution [3]. Additionally, downstream processing of products possibly contaminated with viruses (e.g., human blood derived products and antibodies) requires suitable elimination steps regarding various viruses [3]. Therefore, evaluation and validation of alternative virus purification or virus removal methods such as UF are of interest since UF has advantages over other purification methods including low energy consumption, simplified scale-up, and no chemicals being introduced into the process [1].

Virus removal and/or concentration have been reported for some viruses [2, 4, 5]. The logarithmic titer reduction is in the range of 1 to 8.5 [4] and using UF, oncoretroviral vectors were concentrated 16 to 25 times [2]. Though, at this time no standardized validation protocol for membrane-based virus filtration exists and processing properties are specific to individual virus formulations [3, 4]. Thus, filtration of viruses that are of industrial interest has to be characterized to assess process efficiency and product yield.

AcMNPV has been widely used as foreign gene expression vector and was recently used to develop a vaccine against human papillomavirus [6]. The virus has attracted interest as a vector for gene therapy and is used as a potential insect biopesticide (e.g., as VPN80 [7] for cabbage and cotton cultivation). The rod-shaped budded virions (BV) (DNA genome, enveloped, 40 x 300 nm in size) of AcMNPV, a baculovirus pathogen for lepidopteran insects, are used in cell culture [8]. Previously a study in regard to BV storage at 4 °C, exposure to temperatures from 15 to 65 °C, and to shear stress through stirring and pumping was carried out [3]. The study presented here demonstrates the filtration behavior of a BV solution when filtration parameters with insignificant impact on virus infectivity or particle size are used.

3 Materials and Methods

Production of AcMNPV particles

Cells of the IPLB-SF21-AE (SF-21) cell line (Invitrogen) adapted to Sf-900 II Serum Free Medium (SFM) (pH 6.1; 360 mOsmoles/kg; protein-free; a typical insect medium [9] containing mainly amino acids, glucose, CaCl₂, MgCl₂, KCl and MgSO₄; Invitrogen) were grown at 27°C in SFM supplemented with 5 mg/l gentamicin (Invitrogen) and passaged twice weekly. The virus vHSGFP is a recombinant of AcMNPV that contains the egfp controlled by the Drosophila heat shock protein 70 (hsp70) promoter [10]. vHSGFP was grown in TC-100 medium (Invitrogen) supplemented with 10% FBS (Invitrogen) [3, 8].
SF-21 cells were infected with vHSGFP in suspension culture at a multiplicity of infection (MOI) of 0.1 plaque forming units per cell (pfu/cell) [8]. BV in the supernatant was harvested 4 days post infection (p.i.) by centrifugation (1000 x g, 10 min, at room temperature) and stored at 4 °C (Figure 1) [3, 8].

Ultrafiltration and in-process detection

A commercial ultrafiltration system (Sartoflow Slice 200 Benchtop, Sartorius) was used to concentrate BV solutions. Polyethersulfone (PES) flat sheet membranes (30, 50, 100 or 300 kDa molecular weight cut-off (MWCO), respectively, 0.1 μm pore size (which equates to about 1000 kDa, correlated using electron microscopy [11]); 19.5 x 4.5 cm effective filter area, Sartocon Slice 200, Sartorius) were flushed before usage with 100 ml water (purified with the Q-Gard 2 Purification Pack, Millipore Co.; demineralized; not de-gassed; autoclaved). The trans membrane pressure (TMP) and the trans membrane flux were logged during processing (WinWedge® 32 software, Pressure Sensor Connection Box and TE4101 balance, Sartorius). The flow parallel to the membrane was set to 150 ml/min [5, 12]. The feed vessel was equipped with BV solution (500 ml) and samples (from the feed and permeate each with 1ml) were collected at 100, 200, 300, 400 and 450 ml (± 10 ml) of solution that passed the filter and were stored at 4 °C in opaque containers. All filtration experiments were performed at room temperature (~ 25 °C) and in duplicate. Membranes were cleaned by back-flushing with water from the permeate to the feed (10 min, 500 ml/min), then incubated for > 24 h at 50 °C in sodium hydroxide (1 mol/L, Fisher Scientific) and stored at 4 °C in a solution of sodium hydroxide (0.1 mol/L) with an addition of 20 vol% ethanol (Aaper Alcohol Co.) [5]. The pseudo-steady state flux of water was detected as a benchmark before virus filtration experiments. The PSF was taken when the flux loss was less than 1 ml/min during 5 min or more. Filtered BV solution was reused if no contamination and a viral titer ~ 5 x 10^6 pfu/ml ± 90% was present.

Virus infectivity

TCID_{50} endpoint dilution [8] based on the expression of the Green Fluorescent Protein by host cells infected with vHSGFP and an Eclipse TE200 microscope (green fluorescence emission filter, X200 magnification, Nikon) was utilized to assess the viral titer 5 days p.i.
4 Results and Discussion

The course of infectivity in the retentate during filtration with membranes ranging from 30 to 1000 kDa was determined using end-point dilution (Figure 2).

Figure 2: BV infectivity of retained solution versus the volume passing the membrane during filtration. The filtration system was equipped with 500 ml BV solution, operated at 150 ml/min volume flow parallel to the membrane and was stopped when 450 ml passed the membrane. Virus titer was determined by end-point dilution. Filtrations were done in duplicate (black symbols indicate the first run, repetitions post cleaning are shown using white symbols). Solid lines are added for visualization.

Considering the uncertainty of end-point dilution (0.5 log units [8]), the infectivity of BV solution retained using the 30 kDa membrane was constant. In a half-logarithmic plot the course of infectivity using membranes ranging from 50 to 100 kDa was linear and had a lower slope compared to the use of a higher MWCO. Infectivity increased gradually to about +1 log unit. This might be due to BV partly accumulated in a thin cake layer in front of the membrane. Opposing the 300 and 1000 kDa filters, viral concentration values initially remained constant and then increased linear with a higher slope, slightly higher values of the final concentration and with increased repeatability. Thus, assumed clogging of these membrane pores by BV appears to occur at early times of filtration followed by exponential accumulation of BV in the retentate. High values of viral concentration of about 20 times were archived using 100 to 1000 kDa membranes. This is most likely due to a high ratio of particle average size to membrane pore size [3].

UF using the parameters described reduced infectivity in the range of 2 to 5 log units (Table 1).
Values of the reduction were unordered and point presumably to the interaction between rod-shaped BV particles and the distribution of membrane pore size. Filtration of *Aedes aegypti* densonucleosisvirus (AeDNV) (spherical, 18 – 26 nm in size) decreased infectivity of permeate below the detection limit when a MWCO in the range of 30 to 100 kDa was used. Concurrently, protein concentrations were reported to increase from about 3 to 6 mg/ml in the retentate and to remain constant ~3 mg/ml in solution that passed through the membrane [5, 12]. Since the initial protein concentration was 30 fold higher than it was detected in this study (data not shown), various scenarios including aggregation and different fouling mechanisms, dependent on size and shape of membrane pores and viruses, may explain the retention of AeDNV as opposed by BV that passed membranes ranging from 30 to 100 kDa [13].

### 5 Conclusions

AcMNPV is a baculovirus with broad applications [6, 7], including uses as a protein expression vector. However, the use of viruses for various applications is dependent on its purity and formulation. Membrane based virus filtration is widely used and advantageous in biotechnological and pharmaceutical downstream processing [12, 13], e.g., due to simplified scale-up [1].

In this study, BV of the baculovirus AcMNPV were successfully concentrated 20-fold in a protein-free medium using UF with commercial flat-sheet PES membranes (100 to 1000 kDa MWCO) at an average TMP of 0.15 bar and ~25 °C. Membrane-based filtration appears to be a feasible unit operation in downstream processing of BV for example for industrial scale protein expression. Using membranes and filtration parameters as determined here, our results suggest that reduction of BV in filtered solutions to below detection limits can not be guaranteed using the protocol and the membranes tested here. However,
filtration in the tested MWCO range led to reproducible reduction of BV concentration in the permeate by 2 to 5 log units compared to the unfiltered feed. Hence, membrane-based virus clearance from solutions is beneficial as a unit operation in biotechnological downstream processing since multi-step procedures are generally needed and membrane filtration can significantly reduce the presence of virus as an initial step to facilitate further processing.

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References

