



	Developing a production process for influenza VLPs: a comparison between HEK 293SF and Sf9 production platforms
	Christine M. Thompson, Emma Petiot, Marc G. Aucoin, Olivier Henry, & Amine A. Kamen
Date:	2013
Туре:	Communication de conférence / Conference or Workshop Item
	Thompson, C. M., Petiot, E., Aucoin, M. G., Henry, O., & Kamen, A. A. (2013, June). Developing a production process for influenza VLPs: a comparison between HEK 293SF and Sf9 production platforms [Poster]. 23rd European Society for Animal Cell Technology Meeting (ESACT 2013), Lille, France (2 pages). Published in BMC Proceedings, 7(S6). <u>https://doi.org/10.1186/1753-6561-7-s6-p22</u>

Document en libre accès dans PolyPublie Open Access document in PolyPublie

URL de PolyPublie: PolyPublie URL:	https://publications.polymtl.ca/10677/
Version:	Version officielle de l'éditeur / Published version Révisé par les pairs / Refereed
Conditions d'utilisation: Terms of Use:	CC BY

I Document issued by the official publisher	Document publié chez l'éditeur officiel Document issued by the official publisher	
	Document issued by the official publisher	

Nom de la conférence: 23rd European Society for Animal Cell Technology Meeting (ESACT Conference Name: 2013)

Date et lieu: Date and Location:	2013-06-23 - 2013-06-26, Lille, France
Maison d'édition: Publisher:	BioMed Central
URL officiel: Official URL:	https://doi.org/10.1186/1753-6561-7-s6-p22
Mention légale: Legal notice:	© 2013 Thompson et al.; licensee BioMed Central Ltd. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.

Ce fichier a été téléchargé à partir de PolyPublie, le dépôt institutionnel de Polytechnique Montréal This file has been downloaded from PolyPublie, the institutional repository of Polytechnique Montréal

POSTER PRESENTATION



Open Access

Developing a production process for influenza VLPs: a comparison between HEK 293SF and Sf9 production platforms

Christine M Thompson^{1,2}, Emma Petiot¹, Marc G Aucoin³, Olivier Henry², Amine A Kamen^{1,2*}

From 23rd European Society for Animal Cell Technology (ESACT) Meeting: Better Cells for Better Health Lille, France. 23-26 June 2013

Background

Influenza virus-like particle (VLP) vaccines are one of the most promising approaches to respond to the constant threat of the emergence of pandemic strains, as they possess the potential for higher production capabilities compared to traditional vaccines made in egg-based technology. VLPs are particles produced in cell culture utilizing recombinant protein technology composed of viral antigens that are able to elicit an immune response but lack viral genetic material. Thus far, influenza VLPs have been produced in mammalian, insect and plant based platforms [1], with production in insect cells being the most explored. Baculovirus with mammalian promoters (Bacmam) have been shown to efficiently transduce mammalian cells and further express genes but are unable to replicate, efficiently repressing baculovirus (BV) production that leads to contamination downstream [2]. Influenza VLP production was performed in HEK 293SF cells using the Bacmam gene delivery system. The proposed system was assessed for its ability to produce influenza VLPs composed of Hemagglutinin (HA), Neuraminidase (NA) and Matrix Protein (M1) and compared to VLPs produced in Sf9 cells through the lens of bioprocessing.

Materials and methods

VLPs from both systems were characterized using currently available influenza quantification techniques such as Single Radial Immunodiffusion (SRID) assay, Hemagglutination (HA) assay, Negative Staining Electron Microscopy (NSEM) and western blot.

Results

It was found that VLPs from the HEK 293SF system were present in the culture supernatant in a heterogeneous mixture in terms of particle shape and size. Particles were spherical and also pleomorphic in shape and ranged from sizes of 100-400 nm. Sucrose cushion concentrated samples contained broken particles and a lot of debris. Additionally, it was found that VLPs were associated with the cell pellet after harvest in relatively the same amount as released into the supernatant in the form of unreleased VLPs from NSEM and HA assay analysis. This is possibly due to the sticky nature of the HA protein or from cell clumping during production that worked to trap the VLPs, preventing release into the supernatant. Sf9 cells produced more uniformly shaped VLPs that were spherical in shape, around 100 nm in size and were found to be mainly in the supernatant, not associated with the cell pellet. Sucrose cushion concentrated VLPs contained noticeably less debris than VLPs produced from HEK 293SF cells. It was found that VLP production in Sf9 cells produced 1.5 logs more VLPs than in HEK 293SF cells and had 30× higher HA activity. However, Sf9 VLP samples contained 20× more baculovirus than VLPs, which can contribute to HA activity in both the HA and SRID assays which has to be acknowledged during process development stages. This is the first time to our knowledge that specific production values for influenza VLPs in terms of total particles/ml have been reported.

Conclusions

From this study, the insect-cell baculovirus system produced a more homogeneous population of VLPs compared to its counterpart in HEK 293SF cells. However, this study also highlights the major problem of baculovirus



© 2013 Thompson et al.; licensee BioMed Central Ltd. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.

^{*} Correspondence: amine.kamen@cnrc-nrc.gc.ca

¹Human Health Therapeutics, Vaccine Program, NRC, Montréal, Québec, H4P 2R2, Canada

Full list of author information is available at the end of the article

contamination in the Sf9 system, which requires removal for final vaccine formulations and to help ease the optimization of process production conditions.

Acknowledgements

The authors would like to thank Dr. Ted M Ross of the University of Pittsburgh for kindly donating the Bacmam construct and NSERC for providing the Discovery Grant that supported this study. In addition, we'd like to thank Johnny Montes for his help with viral stock productions and the rest of the ACT group and graduate students at NRC in Montréal for their daily support.

Authors' details

¹Human Health Therapeutics, Vaccine Program, NRC, Montréal, Québec, H4P 2R2, Canada. ²Department of Chemical Engineering, École Polytechnique de Montréal, Montréal, Québec, H3C 3A7, Canada. ³Department of Chemical Engineering, University of Waterloo, Waterloo, Ontario, N2L 3G1, Canada.

Published: 4 December 2013

References

- Kang SM, Song JM, Quan FS, Compans RW: Influenza vaccines based on virus-like particles. Virus research 2009, 143:140-146.
- Tang XC, Lu HR, Ross TM: Baculovirus-produced influenza virus-like particles in mammalian cells protect mice from lethal influenza challenge. Viral immunology 2011, 24:311-319.

doi:10.1186/1753-6561-7-S6-P522

Cite this article as: Thompson *et al.*: **Developing a production process** for influenza VLPs: a comparison between HEK 293SF and Sf9 production platforms. *BMC Proceedings* 2013 7(Suppl 6):P22.

Submit your next manuscript to BioMed Central and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

Submit your manuscript at www.biomedcentral.com/submit

BioMed Central