Poster ID: LB-287

Abstract control N° 10504



Pseudocowpox (PCPV): A next generation viral vector for cancer immunotherapy

A poxviral vector selected for its remarkable ability to induce IFN-alpha

Karola Rittner, Marine Ricordel, Caroline Tosch, Christine Thioudellet, Christine Claudepierre, Virginie Nourtier, Isabelle Farine, Chantal Hoffmann, Doris Schmitt, Benoit Sansas, Johann Foloppe, Philippe Erbs, Nathalie Silvestre, Kaïdre Bendjama, Eric Quéméneur.

AACR 2018 CHICAGO April 14-18, 2018

OBJECTIVE \rightarrow **NEW TOOLS FOR THERAPEUTIC VACCINATION**

Recombinant viruses constitutes a promising modality of immunotherapy. An increasing amount of evidence supports the use of oncolytic viruses in various clinical indications. Also, better understanding of immune mechanisms and progress in immunomonitoring protocols had led to the conclusion that viral induced immune response plays a major role in the anti-tumoral response of this class of viral-based therapeutics. Historically, oncolytic viruses were selected on their ability to lyse tumor cells.

Herein, we describe a next generation anti-cancer virus selected on its capacity to prime and stimulate the immune response. Anti-tumoral efficacy and effects on the immune system were shown in tumor models, by the vector on its own as well as by recombinant virus modified to express relevant tumor antigens.

SCREENING OF *POXVIRIDAE* IN HUMAN PBMCs **★**

We screened a variety of *wt* Poxviridae to identify variants more likely to stimulate the immune response. To this, we exposed human PBMCs to these viruses and measured their capacity to trigger the release of IFN-alpha as a surrogate marker of their pro-immune properties. In parallel, we have screened the impact of virus on viability of PBMC in order to exclude agents with high toxicity for non tumoral cells.

PCPV showed remarkable ability to induce IFN-alpha in human PBMCs.







MVA MVAN33.1 **VV Cop** *wt* Vaccinia Virus Copenhague **CPX** Cowpox **FPV** Fowlpox **PCPV** Pseudocowpox **ORF** Parapoxvirus ovis **MYXV** Myxomavirus **SWPV** Swinepox **YLDV** Yaba-like disease virus **RCN** Raccoonpox **CTV** Cotiavirus

Figure 1: Human Peripheral Blood Mononuclear cells (PBMCs) from two healthy donors per experiment were infected with wt Poxviridae at MOI 0.1, 1 and 5. After o/n incubation, supernatants were isolated and a variety of cytokines and chemokines were quantified (Luminex analysis). The results for IFN-alpha are shown. \rightarrow Among all tested *Poxviridae*, the Parapoxvirus PCPV turned out to be best inducer of IFN-alpha.



Figure 2: Human Peripheral Blood Mononuclear cells (PBMCs) from healthy donors (A, B) were infected with MVA, and the parapoxviruses PCPV and ORF at the MOIs 0.1, 1 and 5. After o/n incubation, cells were stained with LiveDead, the percentage of viable cells is shown. \rightarrow Compared to ORF virus, PCPV showed lower toxicity to PBMCs.

ABOUT PCPV AND THE GENERATION OF ITS RECOMBINANTS

wt Pseudocowpox PCPV

- PCPV strain TJS (ATCC, VR-634) was isolated from a human case of "Milker's nodules", described in Friedman-Kien et al, 1963 Science.
- Patients with Milker's nodules did not develop immunity to vaccinia et vice versa.
- PCPV induces the generation of inclusion and elementary bodies, as it is characteristic for poxviruses.

\rightarrow Recombinant PCPVs

- Recombinant PCPV were generated by insertion of genes into the non-essential VEGF loci by homologous recombination in Bovine Turbinate cells (BT, ATCC CRL-1390).
- Preclinical batches were generated in HeLa cells, crude harvest of infected cells was disrupted using high shear homogenizer clarification by filtration, and purified by tangential flow filtration.
 - PCPV wt
 - **PCPV-mCherry**
 - O PCPV-GFP
 - PCPV-MUC1
 - PCPV-HPV16E7_m



DISCLOSURES

All authors affiliated to Transgene SA are or were employees of Transgene

Transgene SA, Illkirch-Graffenstaden, France

