IFN-I-tolerant oncolytic Semliki Forest virus in combination with anti-PD1 enhances T cell response against mouse glioma

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Oncolytic virotherapy holds promise of effective immunotherapy against otherwise nonresponsive cancers such as glioblastoma. Our previous findings have shown that although oncolytic Semliki Forest virus (SFV) is effective against various mouse glioblastoma models, its therapeutic potency is hampered by type I interferon (IFN-I)-mediated antiviral signaling. In this study, we constructed a novel IFN-I-resistant SFV construct, SFV-AM6, and evaluated its therapeutic potency in vitro, ex vivo, and in vivo in the IFN-I competent mouse GL261 glioma model. In vitro analysis shows that SFV-AM6 causes immunogenic apoptosis in GL261 cells despite high IFN-I signaling. MicroRNA-124 de-targeted SFV-AM6-124T selectively replicates in glioma cells, and it can infect orthotopic GL261 gliomas when administered intra-peritoneally. The combination of SFV-AM6-124T and anti-programmed death 1 (PD1) immunotherapy resulted in increased immune cell infiltration in GL261 gliomas, including an increased tumor-reactive CD8+ fraction. Our results show that SFV-AM6-124T can overcome hurdles of innate anti-viral signaling. Combination therapy with SFV-AM6-124T and anti-PD1 promotes the inflammatory response and improves the immune microenvironment in the GL261 glioma model.

INTRODUCTION

Glioblastoma (GBM) is the most common and most severe form of primary brain tumor in adults.1 The current standard care for GBM consists of radiotherapy, chemotherapy with temozolomide, and surgical resection of the tumor (if possible). None of these interventions is curative, and the median survival remains short (14–15 months) due to relapse in virtually 100% of the cases.2–7

GBM thrives upon the tightly immune-controlled microenvironment in the brain and further skews it toward immunosuppression, making it a challenging target for immunotherapeutic approaches. Recent early phase clinical trials have, however, provided evidence that oncolytic viruses (OVs) can be used for effective GBM immunotherapy in some patients.4–6 Results from preclinical and clinical studies indicate that OV-mediated cancer cell killing and the innate inflammatory response can stimulate an adaptive anti-cancer immune response.7,8 Induction of tumor immunity de novo via virus-induced immunogenic oncolysis should be especially beneficial in treatment of “cold” tumors such as GBM.

Semliki Forest virus (SFV) is an alphavirus (group IV, positive-sense single-stranded RNA [+ssRNA] virus, family Togaviridae), which has a natural capability to penetrate the blood-brain barrier (BBB) in mice. This makes SFV an ideal candidate virus to target brain tumors, provided that viral replication can be restricted to tumor cells. We have previously studied modified oncolytic SFV in different syngeneic mouse glioma model results ranging between encouraging and underwhelming.9–12 The main obstacle in these models is the type I interferon (IFN-I)-mediated innate antiviral response, which inhibits SFV replication in GBM tumors.

IFN-Is play a key role in mediating antiviral immune responses, especially for RNA viruses. Although IFN-I defects are frequent in cancer cells, they cannot be taken for granted in GBM.13,14 Accumulating evidence suggests that IFN-I signaling has major effects on cancer immune surveillance.12 Therefore, despite its negative antiviral effect on OV replication and spread in the tumor tissue, fully functional IFN-I signaling in dendritic cells (DCs) is also needed for the full immunostimulatory therapeutic effect of oncolytic virotherapy. Viruses that can replicate and spread in the cancer cells despite IFN-I signaling would therefore be preferable.

In this study, we aimed to develop a novel oncolytic SFV construct with increased resistance to IFN-I to be used as a potent immunotherapeutic agent against GBM.

RESULTS

SFV-AM6 shows enhanced oncolytic potency in GL261 cells in vitro despite a high IFN-I response

We constructed a novel recombinant SFV termed “SFV-AM6,” which carries 4 aa substitutions as compared to the parental SFV4, that is
nsP3A70G, nsP3Y369F, nsP4R611K and E2K162E (Figure 1A). These mutations were expected to enhance replication and improve in vivo efficacy of the virus based on previous results by us and others (unpublished data).16–18

The plaque phenotype of SFV-AM6 is larger in BHK-21 cells, indicating generally faster replication and spread of the virus (Figure 1B). Notably, SFV-AM6 produces plaques also in GL261 cells, whereas SFV4 does not (Figure 1B). SFV-AM6 showed enhanced GL261 killing potency as quantified using the xCELLigence system (mean ± SD). GL261 cells infected with SFV4 (blue), SFV-AM6 (red) at an MOI of 0.01, or left uninfected (black) (n = 5) are shown. The time point of infection is marked with a black arrowhead. (D) Cell viability of infected GL261 cells was measured with an MTS assay (mean ± SD) 72 h after infection using various MOIs. (E) Caspase-3/7 activity (mean ± SD) in GL261 cells 16 h after infection at an MOI of 1. (F) Virus titers measured (mean ± SD) by plaque titer 24 h after infection at an MOI of 0.01. (G) qRT-PCR analysis of interferon (IFN)-β and indicated IFN-stimulated genes (ISGs) in GL261 cells, 16 h after infection at an MOI of 1 with SFV4 (blue) or SFV-AM6 (red). Data are plotted as mean ± SD. IFIT1, IFN-induced protein with tetratricopeptide repeats 1; TNF, tumor necrosis factor; CXCL10, C-X-C motif chemokine 10; PD-L1, programmed death-ligand 1. Statistical analysis was performed by a Student’s two-tailed, unpaired t test. *p < 0.05, **p < 0.01, ***p < 0.001.

SFV-AM6-infected GL261 cells promote activation and maturation of co-cultured DCs

We have recently published that SFV can induce immunogenic cell death in cancer cells, leading to activation of DCs in vitro,19 and we wanted to analyze whether this occurs also after SFV-AM6 infection in GL261 cells.

Phagocytic activity of mouse immature bone marrow-derived DCs was enhanced when co-cultured with SFV-AM6-infected GL261 cells, as evidenced by increased uptake of pHrodo reagent (Figure 2A) as well as GFP from infected GL261-GFP cells (Figure 2B). Co-culturing with infected GL261 cells also upregulated DC maturation markers CD40, CD80, CD86, and major histocompatibility complex (MHC) class II (Figures 2C–2F). In the case of CD80, CD86, and MHC class
II, upregulation was also observed when DCs were cultured with SFV-AM6 virus alone (MOI of 10), indicating a direct DC activating potency of the virus.

MicroRNA-124 (miR-124) de-targeted SFV-AM6 selectively replicates in GL261 tumors ex vivo and in vivo

To alleviate the neurotoxicity of SFV-AM6 in mice, we constructed a miR-124 de-targeted version of the virus. The design (Figure 3A) was based on our earlier published work. SFV-AM6-124T displayed a large plaque phenotype, similar to SFV-AM6, in BHK-21 and GL261 cells (Figures S2A and S2B; compare with Figure 1), and it displayed SFV-AM6-like oncolytic potency in GL261 cells (Figure S2C), indicating that the viral replication in target tumor cells was not reduced.

To evaluate the infectivity of SFV-AM6-124T in a relevant ex vivo model, we prepared organotypic coronal brain slice cultures from GL261-bearing mice and infected them with SFV-AM6-124T. As seen in Figures 3B and 3C, viral proteins were detected in the GL261 tumor, indicating that SFV-AM6-124T can infect GL261 cells also under ex vivo conditions.

Next, we treated orthotopic GL261 glioma-bearing mice with four intraperitoneal (i.p.) $1 \times 10^7$ plaque-forming unit (PFU) doses of the virus on days 2, 5, 8, and 11 after tumor induction. Immunofluorescence staining revealed tumor regions that were positive both for viral proteins and cleaved caspase-3, 2 days after the last virus injection (Figures 3D and 3E) as well as at the experiment end point (Figures 3F and 3G). Notably, apoptotic tumor areas in SFV-AM6-124T-treated mice were infiltrated by CD11c$^+$ cells (Figures 3F and 3G), which gives evidence of immunogenic cell death in vivo.

To evaluate the therapeutic potential of SFV-AM6-124T, we analyzed the survival of GL261-bearing mice and performed an extensive flow cytometry analysis of immune cell infiltrates in GL261 tumors after therapy with SFV-AM6-124T, anti-PD1, and a combination of both (Figure 4A). SFV-AM6-124T alone did not improve survival of mice (Figure 4B). However, combination therapy improved survival when compared to the control group or virus alone, but it failed when compared to anti-PD1 monotherapy (Figure 4B). Interestingly, flow cytometry analysis did show that the combination therapy significantly increased absolute numbers of all tumor-infiltrating immune cells (CD45$^{high}$) (Figure 4C). Detailed analysis revealed a significant increase in CD3$^+$ T cells (Figure 4D), including CD4$^+$, CD8$^+$, and double-negative (CD3$^-$, CD4$^-$, CD8$^-$) T cell subsets (Figures 4E–4G), B cells (Figure 4H), and DCs (Figure 4H) in the combination therapy. In addition, there was a trend to increase in infiltrating granulocytes, CD11b$^+$ cells, and natural killer (NK) cells in the combination therapy (Figures 4J–4L). In summary, the combination of SFV-AM6-124T and PD1 therapies synergize to improve intra-tumoral immune cell infiltration and inflammation.

SFV-AM6-124T promotes a distinct CD8$^+$ T cell phenotype in the GL261 microenvironment

Encouraged by the significant increase in infiltration of cytotoxic T cells after combination therapy, we characterized them in detail by flow cytometry (Table S2). SFV-AM6-124T monotherapy modestly upregulated the activation markers CD69 and CD25 and the proliferation marker Ki67, but it led to a more distinct upregulation of CD127 and the senescence marker KLRG1 on tumor-infiltrating CD8$^+$ T cells as compared to controls (Figure 5A), as
reflected in higher numbers of KLRG1+CD127- and KLRG1+CD127+CD8+ T cell populations in the virus treatment groups (Figures 5B–5D). Anti-PD1 as monotherapy induced strong expression of Ki67 and activation markers CD69, CD25, CD44, and the homing receptor CD62L on tumor-infiltrating CD8+ T cells, suggesting an effector memory phenotype (Figure 5A). In addition, only anti-PD1 monotherapy induced upregulation of CX3CR1 (Figure 5A; Figures S3A–S3C), a subset of CD8+ T cells known to be responsive to PD1 therapy. After combination therapy, tumor-infiltrating CD8+ T cells had an activated (CD69+CD25+), effector phenotype (CD44+CD62L-) but were in a state of low proliferation (Ki67low) (Figure 5A; Figures S3D–S3F). Combination therapy also markedly increased co-expression of multiple T cell activation/exhaustion markers PD1, LAG3, and TIM3 (Figures 5A and 5E). In addition, there was an increase in intra-tumoral regulatory T cells (Tregs) (CD4+CD25+FOXP3+CD127-) in the combination treatment group (Figures S3G and S3H). The above results indicate that SFV-AM6-124T treatment induced a distinct effector CD8+ T cell phenotype with high KLRG1 expression, and when combined with anti-PD1, it was associated with elevated PD1 and TIM3 expression.

We next analyzed CD8+ T cells targeting GL261 tumor, which is of importance in respect to cancer therapy. The GL261 cells overexpress endogenous retrovirus-based tumor antigens, and T cells against this endogenous antigen are detectable using tetramers against the murine leukemia virus (MuLV) env2-env epitope (KSPWFTTL). We observed an increased infiltration of KSPWFTTL-tetramer (Tet)+ tumor-specific T cells in anti-PD1 monotherapy and in combination therapy compared to controls (Figures 5F and 5G). More importantly, there was no difference in the percentage of KLRG1+CD127+ anti-tumor CD8+ T cells (Figure 5H) and lower levels of terminally differentiated KLRG1+CD127- anti-tumor CD8+ T cells (Figure 5I) in anti-PD1 and combination therapy compared to SFV-AM6-124T monotherapy. In addition, there were no differences in the co-expression of multiple T cell activation/exhaustion markers PD1, LAG3, and TIM3 (Figure 5J) among the different treatment groups. In conclusion, our therapies induced anti-tumor-specific CD8+ T cells, and especially combination therapy rescued the terminally differentiated KLRG1+ T cell phenotype.

DISCUSSION
Given the important role of IFN-I in promoting cancer immunity and evidence of functional IFN-I signaling in human GBM samples, it seems likely that IFN-I resistant/tolerant viruses would be favorable as oncolytic immunotherapy agents. Consistent with this, the clinically relevant poliovirus PVS-RIPO has recently been reported to be tolerant to IFN-I in cancer cells. In the present study, we report a novel SFV-AM6 virus that shows enhanced replication and cytotoxicity in a highly IFN-I-competent GL261 mouse glioma cell line (Figure 1). Importantly, the increase in replication was not due to a dampened IFN-I response (Figure 1G), indicating a clearly improved, IFN-I tolerant phenotype of the SFV-AM6 over wild-type SFV4. A plaque assay in particular gave a strong indication of its ability to spread in restrictive tumor microenvironment-like conditions (Figure 1B). In contrast to regular non-immunogenic apoptosis, SFV-AM6 infection triggers DC activation (Figure 2). This is in line with our previously reported findings of SFV-induced pathogen- and damage-associated molecular pattern (PAMP and DAMP)-driven immunogenic cell death, and it indicates the immunotherapeutic potential of the SFV-AM6 construct.
While the molecular mechanistic explanation for the increased oncolytic potency of SFV-AM6 is unclear, some reasonable speculations can be made. The NsP3A70G substitution is located in the conserved nsP3 macrodomain and is thus possibly linked to viral counteraction of IFN-stimulated poly(ADP-ribose) polymerases.16,17 The NsP3Y369F mutation has been recently shown to inhibit viral hyperactivation of the phosphatidylinositol 3-kinase (PI3K-AKT-mTOR) pathway and reduce replication in vivo.25 The PI3K-AKT pathway is, however, overactivated in many cancers, including GBM, due to frequent loss of the negative regulator PTEN.26 The NsP3Y369F mutation could therefore selectively increase virus replication in cancer cells with oncogenic over-activation in the PI3K-AKT pathway, such as PTEN-defective GL261 cells.27

The NsP4R611K substitution affects the penultimate residue of nsP4 and the overlapping 26S sub-genomic RNA promoter region, and most probably acts by enhancing replication speed and/or promoting synthesis of 26S RNA (unpublished data). Finally, a Lys to Glu switch
at residue 162 in E2 was included in our construct due to its positive effect on induction of viremia as well as BBB penetration.18

miR-124 de-targeting11,20 did not interfere with GL261 infectivity in vitro or ex vivo (Figure S2; Figure 3B and 3C). Our previously published work indicates that although miR-124 de-targeting significantly attenuates neurovirulence, it does not result in complete protection when the virus is administered intracranially.20 Because of this, as well as the advantages of systemic delivery, we decided to study SFV-AM6-124T as an i.p. administered oncolytic agent. In

Figure 5. SFV-AM6-124T promotes a distinct CD8\(^+\) T cell phenotype in GL261 gliomas

(A) MFI heatmap of Ki67, CD69, CD25, CD44, CD62L, CD127, CX3CR1, KLRG1, PD1, TIM3, and LAG3 expression in CD8\(^+\) T cells from control and treated tumors. (B) Flow cytometry gating strategy for quantification of KLRG1 and CD127-expressing CD8\(^+\) T cells. (C and D) Quantification of KLRG1\(^+\)/CD127\(^-\) (C) and KLRG1\(^+\)/CD127\(^+\) (D) cells. (E) Quantification of PD-1, TIM3, and LAG3 co-expression on CD8 T cells with SPICE software. Colors of the pie arcs depict the expression of individual inhibitory receptors, while the pie depicts the average proportion of co-expressed inhibitory receptors. (F) Bivariate plots showing staining for tetramer (Kb-restricted peptide amino acids 604–611 of p15E protein [KSPWFTTL]) on tumor-infiltrating CD8 T cells, and (G) the quantification of tumor-infiltrating tetramer\(^+\) CD8 T cells in each treatment group. (H and I) Quantification of KLRG1\(^+\)/CD127\(^+\) (H) and KLRG1\(^+\)/CD127\(^+\) (I) tetramer\(^+\) CD8 T cell subpopulation. (J) Quantification of PD-1, TIM3, and LAG3 co-expression on tetramer\(^+\) CD8 T cells with SPICE software. Panels C, D, G, H and I are plotted as mean ± SD. Statistical analysis was done with one-way ANOVA with Tukey’s post hoc test (C, D, H and I) or Kruskal-Wallis with Dunn’s post hoc test (G). *p < 0.05, **p < 0.01, ***p < 0.001.
in the case of SFV, we consider also that infection/activation of brain endothelial cells through systemic delivery should favorably increase immune cell infiltration. Impressively, systemically administered SFV-AM6-124T can reach orthotopic GL261 tumors, resulting in apoptotic/dead regions surrounded by SFV-infected cells (Figures 3D and 3E). The infected regions were infiltrated by CD11c+ cells (Figures 3F and 3G), which are likely peripheral DCs and serve as an indication of in vivo immunogenic oncolysis. Evidence of immunogenic oncolysis has been reported previously in the GL261 model with Newcastle disease virus\textsuperscript{29} as well as in a clinical trial with parvovirus (ParvOryx).\textsuperscript{29}

In the context of immunotherapy, we analyzed the effect of SFV-AM6-124T to the immune microenvironment in GL261 gliomas both as monotherapy and when combined with an anti-PD1 antibody, as anti-PD1 therapy has been shown to be effective against GL261 gliomas.\textsuperscript{30} Anti-PD1 combination was considered rational also due to SFV-induced PD-L1 expression in GL261 cells (Figure 1G), likely via IFN-I signaling.\textsuperscript{31,32} In the present study, combination therapy with virus and anti-PD1 improved survival when compared to the control group or virus alone, but it failed to improve survival when compared to anti-PD1 alone (Figure 4A). The combination therapy group actually showed somewhat reduced survival when compared to anti-PD1. The lack of statistical significance between these two groups can be explained by the lower number of mice in the anti-PD1 group. Despite lacking clear survival benefit, the combination therapy, however, increased general numbers of GL261-infiltrating T cells, B cells, DCs (Figure 4), and intra-tumoral inflammation, suggesting potential pseudo-progression of the tumors, a phenomenon frequently observed after OV therapy of cancers.\textsuperscript{33}

Detailed analysis of the tumor-infiltrating cytotoxic CD8\textsuperscript{+} T cells revealed that SFV-AM6-124T promoted a low-proliferating and exhausted KLRG1\textsuperscript{+} effector phenotype (Figure 5), which is likely associated with the cumulative T cell receptor and inflammatory stimulus.\textsuperscript{34} Although KLRG1\textsuperscript{+}CD8\textsuperscript{+} cells have been shown to differentiate into “exKLRG1” cells (KLRG1\textsuperscript{−}CD127\textsuperscript{−}) with potent antiviral and anti-tumor effects,\textsuperscript{34} it is reasonable to speculate that their presence is a therapeutically unfavorable response. The KLRG1\textsuperscript{+}CD127\textsuperscript{−} population was also increased in the GL261-targeting CD8\textsuperscript{+} T cells in response to virus alone (Figure 5I). The combination with anti-PD1, however, reduced the population to anti-PD1 monotherapy levels (Figure 5I), indicating reversion of the SFV-AM6-124T-driven negative effect in anti-tumor T cells.

It is not self-evident that OVs would synergize positively with an anti-PD1-boosted T cell response. In fact, it can be expected that the immunodominant viral antigens could also redirect the T cell response away from anti-tumor toward anti-virus and promote T cell exhaustion. A virus-induced IFN-I response can also negatively affect anti-tumor T cells, as reported recently by Evgin et al.\textsuperscript{35} using a combination of oncolytic vesicular stomatitis virus (VSV) and chimeric antigen receptor (CAR) T cell therapy in a mouse melanoma model. Herein, we report that systemically administered oncolytic SFV-AM6-124T in combination with anti-PD1 results in increased anti-tumor CD8\textsuperscript{+} infiltration in an orthotopic mouse GL261 model. Importantly, the GL261-reactive CD8\textsuperscript{+} T cells did not show a similar terminally differentiated phenotype that was observed in the total CD8\textsuperscript{+} population (Figure 5), indicating that the anti-tumor response is not negatively affected by the virus.

The lack of correlation to survival benefit in SFV-AM6-124T-treated mice makes interpretation of the results challenging. It is likely that the kinetics of the combination therapy-induced anti-tumor response is not sufficiently fast to prevent tumor growth within the short time frame of the GL261 model. The symptoms observed in the combination therapy group could be due to heavily increased inflammation in the tumors. Although some level of inflammation in the intracranial tumors is a favorable immunotherapeutic response, it can lead to severe symptoms in mice. We therefore advocate that monitoring mouse symptoms is not sufficient, and analysis of the immune cell infiltrates should be a key consideration when evaluating the therapeutic potency.

Taken together, our results support that the oncolytic potency of SFV-AM6-124T can be harnessed in combination with anti-PD1 to enhance the anti-tumor immune response. The results can be translated into development of effective oncolytic immunotherapy against GBM.

MATERIALS AND METHODS

Cell lines

GL261 mouse GBM cells (gift from Dr. G. Sáfrány, National Research Institute for Radiobiology and Radiohygiene, Budapest, Hungary) were cultured in Gibco DMEM GlutaMAX (Thermo Fischer Scientific, Waltham, MA, USA) supplemented with 10% Gibco heat-inactivated fetal bovine serum (FBS, Thermo Fischer Scientific), 10 U/mL Gibco penicillin-streptomycin (PEST, Thermo Fisher Scientific), and 1 mM Gibco sodium pyruvate (Thermo Fisher Scientific). Baby hamster kidney (BHK)-21 cells were cultured in Gibco Glasgow’s minimum essential medium (GMEM, Thermo Fisher Scientific) supplemented with 10% FBS, 2% tryptose phosphate broth (Teknova, Hollister, CA, USA), 20 mM HEPES, and 10 U/mL Gibco PEST (Thermo Fisher Scientific).

Viruses

All modifications/mutations were made into pCMV-SFV4 backbone plasmid\textsuperscript{36} either by \textit{in vitro} mutagenesis and/or by replacement of fragments with the mutation-containing fragments from previously made constructs; the obtained construct was verified by sequencing and designated as pCMV-SFV-AM6. pCMV-SFV-AM6-124T was constructed by cloning the miR124T fragment from pSP6-SFV4-124T\textsuperscript{37} into pCMV-SFV-AM6. All SFV viruses were produced and titrated in BHK-21 cells as described in Supplemental materials and methods.

GL261 infection assays

The xCELLigence system (ACEA Biosciences, San Diego, CA, USA) was used to assess GL261 de-attachment following an MOI of 0.01
virus infection of 10,000 cells. The assay was performed as described in Supplemental materials and methods. For the cell viability assay, GL261 cells were seeded onto a 96-well plate (20,000 cells/well) and infected on the next day with virus (diluted in complete cell culture medium) at different MOIs. Cell viability was measured using a Cell-Titer 96 AQueous One solution cell proliferation assay (Promega, Madison, WI, USA) according to the manufacturer’s instructions 72 h after infection. A Caspase-Glo 3/7 assay system (Promega) was used to measure caspase-3 and caspase-7 activity in GL261 cells (plated onto a 96-well plate as above) infected with an MOI of 1, 16 h after infection. A plaque assay in GL261 cells was performed according to a titration protocol (Supplemental materials and methods).

**Infection of brain slice culture**

Brains from GL261 tumor-bearing mice were collected in complete Hanks’ balanced salt solution (HBSS; 2.5 mM HEPES [pH 7.4], 30 mM d-glucose, 1 mM CaCl₂, 1 mM MgSO₄, 4 mM NaHCO₃) and embedded in low-melting point agarose on ice. Agarose-slices were transferred onto the bottom of wells. Slice cultures were incubated (37°C, 5% CO₂) for 1 h, after which 5 μL of added culture medium (68% DMEM + L-glutamine, 200 mM NaHCO₃, 30 mM D-glucose, 1 mM CaCl₂, 1 mM MgSO₄, 4 mM NaHCO₃) at different MOIs. Cell viability was measured using a CellTiter 96 AQueous One solution cell proliferation assay (Promega, Madison, WI, USA) according to the manufacturer’s instructions 72 h after infection. A Caspase-Glo 3/7 assay system (Promega) was used to measure caspase-3 and caspase-7 activity in GL261 cells (plated onto a 96-well plate as above) infected with an MOI of 1, 16 h after infection. A plaque assay in GL261 cells was performed according to a titration protocol (Supplemental materials and methods).

**qRT-PCR**

500,000 GL261 cells were infected at an MOI of 1. Infected and uninfected cells (three parallel samples each) were harvested 16 h after infection. RNA was extracted using an RNeasy mini kit (QIAGEN, Germany). cDNA was synthesized from pooled RNA samples with iScript cDNA synthesis (Bio-Rad, Hercules, CA, USA) from 1 μg of extracted RNA. iQ SYBR Green supermix (Bio-Rad) and the primers listed in Table S1. Each sample was run in duplicate, and the fold change of expression was quantified using the 2^–ΔΔCt method.

**Flow cytometry**

Single cells from the murine brain earing GL261 tumors were isolated using a mouse tumor dissociation kit (Miltenyi Biotec, Germany) and a gentleMACS Octo dissociator with heaters (Miltenyi Biotec). Myelin was depleted using 25% BSA density gradient centrifugation for 20 min at 2,600 rpm and subsequently removing the myelin rim on top of the BSA gradient. CD45⁺ cells were enriched using mouse CD45 MicroBeads (Miltenyi Biotec) following the manufacturer’s instructions. Unspecific Fc receptor binding in all single-cell suspensions was then blocked by using anti-mouse CD16/CD32 antibody (clone 93, BioLegend, San Diego, CA, USA), and were then stained using fluorochrome-conjugated antibodies (Table S2). All antibodies were diluted 1:100 from stock concentration in Brilliant Violet stain particles conjugate for phagocytosis (Thermo Fisher Scientific) and Alexa Fluor 647 (AF647)-conjugated goat anti-hamster (Thermo Fisher Scientific) used as secondary antibodies. The sections were imaged with an Eclipse Ti-S microscope (Nikon, Japan).

**DC phagocytosis and maturation assays**

Bone marrow-derived DCs were isolated from the femur and tibia of female 7- to 8-week-old wild-type C57BL/6N (Janvier Labs, France) and cultured in Gibco Iscove’s modified Dulbecco’s medium (IMDM, Thermo Fisher Scientific) supplemented with 10% heat-inactivated FBS, 10 U/mL PEST, 10 mM HEPES, 50 mM β-mercaptoethanol, 20 ng/mL recombinant murine interleukin (IL)-4, and 20 ng/mL recombinant murine granulocyte-macrophage colony-stimulating factor (GM-CSF, Nordic BioSite, Sweden). The medium was changed every 3 days. Non-adherent immature DCs were harvested for a co-culture assay on day 7.

For measuring phagocytic activity, GL261 cells were infected with SFV-AM6 at an MOI of 10 for 48 h followed by co-culturing with DGs (1:1 ratio) and 100 μg of pHrodo Red Staphylococcus aureus Bio-particles conjugate for phagocytosis (Thermo Fisher Scientific) for 2
h. To measure phagocytosis of tumor cells directly, GL261-GFP cells were infected and co-cultured with DCs as above. Noninfected cells were used as a control. Cells were harvested and DCs were stained with anti-CD11c-allophycocyanin (APC)/Cy7 (BioLegend). Phagocytosis of pHrodo reagent was quantified as the percentage of CD11c+GFP+ cells. Similarly, the phagocytosis of dead GL261-GFP cells by DCs was quantified as the percentage of CD11c+GFP+ cells

For the DC maturation assay, GL261 cells were infected with SFV-AM6 (MOI of 10) for 48 h followed by co-culturing with immature DCs at a ratio 1:1 for 18 h. Noninfected GL261 cells were used as a negative control. Additionally, SFV-AM6 was added directly on DCs (MOI of 10). Maturation of DCs was analyzed by flow cytometry using anti-CD11c-APC/Cy7, anti-CD11b-peridinin chlorophyll protein (PerCP), anti-CD80-Pacific Blue, anti-CD86-phycocerythrin (PE)/Cy7, anti-CD40-APC, and anti-I-A/I-E-Brilliant Violet 510 (BV510, BD Biosciences) antibodies.

Statistical analysis
All statistical analysis was performed using GraphPad Prism software.

Ethics statement
The Swedish Work Environment Authority has approved the work with genetic modification of SFV (ID no. 202100-2932 v66a14 [laboratory] and v67a10 [mice]). All experiments regarding modified SFV were conducted under biosafety level 2. The local Animal Ethics Committee in Stockholm (N164/15, N170/13) approved the animal studies.

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.omto.2021.03.008.

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AUTHOR CONTRIBUTIONS
M.M., M.R., R.L., A.D., D.Y., A.M., and M.E. designed the experiments. M.M., M.R., R.L., J.M., M.-M.M., and A.M. conducted the experiments. M.M., M.R., R.L., J.M., A.D., D.Y., A.M., and M.E. wrote the paper.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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