Oncolytic poxvirus CF33-hNIS-ΔF14.5 favorably modulates tumor immune microenvironment and works synergistically with anti-PD-L1 antibody in a triple-negative breast cancer model

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ABSTRACT

Triple-negative breast cancer is the most aggressive subtype of breast cancer and is difficult to treat. Breast cancer is considered to be poorly immunogenic and hence is less responsive to immunotherapies. We tested whether the oncolytic poxvirus CF33-hNIS-ΔF14.5 could modulate tumor immune microenvironment and make the tumors responsive to the immune checkpoint inhibitor anti-PD-L1. We found that virus infection causes the upregulation of PD-L1 levels on triple-negative breast cancer cells in vitro as well as in vivo in mice. In a mouse model of orthotopic triple-negative breast cancer, the virus was found to increase tumor infiltration by CD8+ T cells. Likewise, in mice treated with CF33-hNIS-ΔF14.5 high levels of proinflammatory cytokines IFNγ and IL-6 were found in the tumors but not in the serum. The levels of immune modulation were even higher in mice that were treated with a combination of the virus and anti-PD-L1 antibody. While CF33-hNIS-ΔF14.5 and anti-PD-L1 antibody failed to exert significant anti-tumor effect as a single agent, a combination of the two agents resulted in significant anti-tumor effect with 50% mice experiencing complete tumor regression when both agents were injected intra-tumorally. Furthermore, the ‘cured’ mice did not develop tumor after re-challenge with the same cancer cells suggesting that they developed immunity against those cancer cells. Taken together, our study shows that CF33-hNIS-ΔF14.5 favorably modulates tumor immune microenvironment in triple-negative breast cancer model making them responsive to the immune checkpoint inhibitor anti-PD-L1, and hence warrants further studies to determine the clinical applicability of this combination therapy.

Introduction

Breast cancer (BrCa) is the most common type of cancer in women and the leading cause of cancer-related death in women worldwide. Surgery and radiation therapy are commonly used for the treatment of primary BrCa, while chemotherapy and hormone therapies are standard for the management of metastatic BrCa. Treatment of this highly heterogeneous disease is challenging because of differential responses of BrCa subtypes and subpopulations to these therapies. Triple-negative BrCa (TNBC), a subtype of BrCa lacking estrogen receptor, progesterone receptor and human epidermal growth factor receptor (HER2), has the poorest prognosis amongst all subtypes. Despite an improvement in diagnosis and treatment, 5-year survival for metastatic BrCa is less than 30% and almost all patients with TNBC succumb to their disease, therefore alternative therapeutics with better efficacy are urgently needed.

In the last 2 decades, the field of immunotherapy has seen major breakthroughs that have established immunotherapy as a major therapeutic approach for cancer. In particular, immune checkpoint inhibitors (ICIs) targeting CTLA-4 and PD1/PD-L1 have shown unprecedented long-lasting therapeutic efficacy in different types of malignancies. However, despite these unprecedented long-lasting response rates, only a small fraction of patients benefit from ICIs. Anti-tumor efficacies of ICIs are mostly limited to tumors with T cell-inflamed tumor microenvironment (TME) also known as ‘immunologically hot’ tumors. Poorly immunogenic tumors with scarce T cells, also referred to as ‘immunologically cold’ tumors, are largely refractory to ICIs. Hence, a combination of ICI with other therapeutics that could potentially convert immunologically ‘cold’ TME to ‘hot’ may allow to harness the benefits of ICIs in a broader patient population.

Oncolytic viruses (OVs) are naturally occurring or genetically modified viruses that can selectively replicate in and kill cancer cells while sparing nonmalignant cells. In addition to direct cell killing by the virtue of cancer-selective replication, OVs exert indirect anti-neoplastic effect through the destruction of tumor vasculature as well as activation of innate and adaptive immune system. Traditionally, the field of oncolytic virotherapy was focused mostly on improving the replication potential of OVs based on the concept that direct killing of cancer cells by OVs is the main mechanism of action. However, recent studies have shown that the success of OVs is as much as, or more, dependent on their immune-
modulating capability as it is on their ability to directly kill cancer cells.\textsuperscript{15-19} OVs could modulate the immunological landscape in TME through a variety of different mechanisms.\textsuperscript{20} For example, OVs have been shown to induce immunogenic cell death which plays an important role in the induction of adaptive immunity.\textsuperscript{21-23} Furthermore, OVs can modulate cytokine/chemokine in TME leading to increased tumor infiltration by CD8\(^+\) T cells and other immune cells.\textsuperscript{9} However, CD8\(^+\) T cells within the TME may be blocked by cancer cells through checkpoint proteins such as PD-L1 in which case addition of ICIs should be helpful. Taken together, OVs have the potential to set the stage for ICIs and ICIs have the ability to allow an unperturbed activity of OV-activated anti-tumor immunity; hence, it is logical to surmise that OVs and ICIs may result in synergistic anti-tumor effect.

CF33-hNIS-ΔF14.5 is a chimeric poxvirus which has demonstrated strong oncolytic effect against several tumor models at relatively low doses.\textsuperscript{21,24-26} We have previously reported that CF33-hNIS-ΔF14.5 induces immunogenic cell death and increases CD8\(^+\) T cells infiltration in tumors.\textsuperscript{21} Here, we studied the therapeutic potential of CF33-hNIS-ΔF14.5 in combination with an anti-PD-L1 (α-PD-L1) antibody in a triple-negative breast cancer model.

Materials and methods

Cell lines

Human triple-negative breast cancer cells MDA-MB-468, Hs578T, and murine breast cancer cells 4T1 were purchased from American Type Culture Collection (Manassas, USA). The murine triple-negative breast cancer cells E0771 were a kind gift from Stephen J. Forman’s laboratory at City of Hope, Duarte, USA. All cells were cultured in RPMI-1640 medium supplemented with 10% fetal bovine serum, 2 mM L-glutamine and 100 U/ml penicillin-streptomycin. Cells were maintained in a humidified incubator at 5% CO\(_2\) and 37°C.

Virus proliferation and cytotoxicity assays

In order to determine the growth kinetics of the virus, cancer cells were seeded in 6 well plates at the density of 300000/well in 2 mL cell culture medium. Next day, cells were infected with the virus at multiplicity of infection (MOI) 0.03. Cell lysates were collected at 24-, 48- and 72-h post-infection and virus titer in the lysates were determined by standard plaque assay on CV1 cells as described previously.\textsuperscript{21}

To determine the cytotoxic potential of the virus, 3000 cells were seeded per well of 96-well culture plates in 100 μl cell culture medium. The following day, cells were either mock-infected or infected by CF33-hNIS-ΔF14.5 at MOIs 0.01, 0.1 or 1. After infection, plates were incubated for 72 h. Next, CellTiter96\textsuperscript{A}Queous (Promega) reagent was added to the cells and 1 h later absorbance was measured at 490 nm using a plate reader (Tecan Spark) as per the manufacturer’s instruction. Survival was calculated relative to mock-infected wells.

Virus and therapeutic αPD-L1 antibody

Construction of CF33 has been described previously.\textsuperscript{21} The virus CF33-hNIS-ΔF14.5 is deleted of the genes J2R and F14.5L and it encodes human sodium iodide symporter (hNIS) gene. CF33-hNIS-ΔF14.5 was amplified in CV1 cells and purified on sucrose gradients. Mouse-specific α-PD-L1 antibody (Cat#BP0101; Clone 10E9.9G2) was purchased from Bio X Cell (West Lebanon, NH, USA). We chose to use this antibody as Liu \textit{et al.} have previously shown that this antibody works well in C57BL/6 mice.\textsuperscript{27}

Tumor model and treatment

All animal studies were conducted under City of Hope Institutional Animal Care and Use Committee approved protocol (IACUC#15003) in compliance with NIH’s guideline for the use of laboratory animals.

Female C57BL/6 mice, 4–6 weeks old, were purchased from Charles River and acclimatized for 1 week in a pathogen-free environment at City of Hope’s animal facility. To generate orthotopic tumors, E0771 cells were used. A total of 10\(^5\) cells in 50 μL volume (PBS and matrilgel 1:1) were injected in the abdominal mammary fat pad. One tumor per mouse was generated for tumor regression and survival studies whereas two tumors (bilateral orthotopic tumors) per mouse were generated for studies involving immune analysis. For a different tumor regression and survival study, bilateral tumors were generated by injecting 10\(^5\) E0771 cells in right-side mammary fat pad and 2 × 10\(^4\) E0771 cells in left-side mammary fat pad. When tumors became palpable, mice were sorted into different treatment groups such that the average tumor volume in each group was similar. Mice were injected intra-tumorally with 10\(^7\) PFU of CF33-hNIS-ΔF14.5 or 100 μg of α-PD-L1 either intra-tumorally or intra-peritoneally, as shown in the treatment schemes of respective figures, on each of experimental days 1, 3 and 5. Mice were weighed twice weekly and tumor volumes were measured twice weekly using digital calipers. During the entire experimental period, no overt toxicity was observed in any mouse and all mice continued to gain weight. Tumor volumes were calculated as described previously and mice were euthanized when tumors exceeded 2500 mm\(^3\) in volume.\textsuperscript{21} For tumor re-challenge, mice that were tumor free for 30 days were injected with 10\(^5\) E0771 cells in the abdominal mammary fat pad on the opposite side from the regressed tumors. Age-matched naïve mice were used as a control for the re-challenge study.

For immune analysis, mice bearing bilateral tumors were euthanized on day 7 after the first dose of treatment and their tumors and serum were harvested. One tumor was collected in PBS for flowcytometric analysis and the other tumor was divided into two halves. One half was weighed and snap-frozen in liquid nitrogen to be later used for cytokine analysis and the second half of tumors were
collected in formalin for immunohistochemical (IHC) analysis. For FACS, harvested tumors were weighed, and single cells were generated using mouse tumor dissociation kit (Miltenyi Biotec; Cat# 130-096-730) and the GentleMACS™ dissociator (Miltenyi Biotec). Single cells were frozen in 90% FBS and 10% DMSO and stored at −80°C before staining for flow cytometric analysis.

**Flow cytometry**

Cells were infected with CF33-hNIS-ΔF14.5 at an MOI 3 or were mock-infected. Eighteen hours post-infection, cells were detached from the plates using 5 mM EDTA. Cells were washed twice with PBS and stained with α-PD-L1 antibody or isotype antibody. Cells were analyzed on BD Accuri™ C6 cytometer.

For analysis of immune cells in E0771 tumors, single cells from tumors that were frozen in 90% FBS and 10% DMSO and stored at −80°C were thawed. Cells were washed twice with PBS and stained with zombie UV fixable viability dye (BioLegend; Cat# 423108) for 30 min in PBS. Cells were fixed with 4% paraformaldehyde for 15 min and then stained with PerCP/Cy5.5-conjugated CD45 antibody (BioLegend; Cat# 103131, Clone: 30-F11), VioGreen-conjugated CD8 antibody (Miltenyi Biotec; Cat# 130-109-252, Clone: REA601), FITC-conjugated CD4 antibody (BioLegend; Cat#130308, Clone: H129.19) and PE-conjugated PD-L1 antibody (BD Biosciences; Cat#558091, Clone: MIH5) for 30 min at 4°C in dark. In a separate staining panel, cells were stained for surface antigens and permeabilized using BD fix/perm kit (BD Biosciences; Cat#554714). After permeabilization cells were stained with APC-conjugated IFNγ antibody (BioLegend; Cat#505810, Clone: XMG1.2). After staining, cells were fixed with 4% paraformaldehyde for 15 min and data collected on BD LSRFortessa.

**Cytokine measurement**

Tumors were homogenized in lysis buffer (10 mM Tris-HCl pH 8.0, 150 mM NaCl, 1% NP-40, 10% glycerol, 5mM EDTA and 1x protease inhibitors). Tumors were lysed in volumes 5 times their weight, for example, 200 μg of tumor was lysed in a total volume of 1 mL lysis buffer. Lysates were briefly sonicated in a water bath sonicator and spun down at 14000 g for 15 min at 4°C. Supernatants were transferred to new tubes and 25 μL from each sample was used to determine cytokines concentration using LEGENDplex Mouse Th CytoKine Panel kit (Cat#740741; BioLegend) following manufacturer’s instruction. Likewise, blood was collected by cardiac puncture at the time of euthanizing the mice. Serum was collected from the blood samples and 2 fold diluted serum was used to determine cytokines concentration using LEGENDplex Mouse Th CytoKine Panel kit (Cat#740741; BioLegend) following manufacturer’s instruction. Samples were run in duplicate. BD Accuri™ C6 flowcytometer was used to acquire data and data analysis was performed using LegendPlex software.

**Immunohistochemical analysis**

Tumors were fixed in formalin for 48 h, embedded in paraffin and sliced into 5 μm thin sections. Prior to staining, tumor sections were deparaffinized and rehydrated, after which heat-mediated antigen retrieval was performed as previously described. Endogenous peroxidase was quenched by incubating the slides in 10% freshly prepared H2O2 for 10 min. Sections were blocked with TMB blocking buffer (PerkinElmer; Cat#FP1020), and then incubated with rabbit anti-CD8 antibody (Abcam; Cat#ab209775, Clone: EPR20305) or rabbit anti-CD4 antibody (Abcam; Cat#ab183685, Clone: EPR19514) or rabbit anti-PD-L1 antibody (Cell Signaling Technology; Cat#64988, Clone: D5V3B) overnight at 4°C. The following day, sections were washed and treated for 1 h at room temperature with HRP-conjugated polyclonal goat anti-rabbit antibody (Abcam; Cat#ab6721). Next, sections were washed, and brown color was developed using DAB substrate kit (Abcam; Cat#ab64238) following the manufacturer’s protocol. Finally, slides were counterstained with hematoxylin (Sigma-Aldrich; MHS16-500), sections were dehydrated, and coverslips were applied using permount (Fisher Scientific; Cat#SP15-100) as a mounting medium.

Stained tumor sections were imaged using the whole slide imager NanoZoomer S360 (Hamamatsu, Japan). Images were analyzed and quantified using QuPath software. Necrotic areas were excluded for quantification purpose and data are presented as positive cells/mm² of tumor section.

**Statistical analysis**

For comparing the means of more than two groups one-way ANOVA was used with a 95% confidence interval. P-values <0.05 were deemed significant. Tumor growth curve (average tumor volume) for different treatment groups was compared using two-way ANOVA with Dunnett’s test. Survival studies were analyzed for statistical significance using the
log-rank Mantel–Cox test. Studies involving tumor regression and survival in mice were performed on $n = 7–8$ mice per treatment group so as to obtain reasonable statistical value. GraphPad Prism 5 Software (GraphPad Software, La Jolla, CA, USA) was used to calculate statistical values.

**Results**

**Breast cancer cells upregulate PD-L1 expression after infection with CF33-hNIS-ΔF14.5**

In order to investigate the effect of virus infection on PD-L1 expression in triple-negative breast cancer, we infected human TNBC (MDA-MB-468 and Hs578T) and murine TNBC cells (4T1 and E0771) with CF33-hNIS-ΔF14.5 at MOI 3 or mock-infected. Eighteen hours post-infection cells were analyzed for PD-L1 levels on cell surface. Basal levels of PD-L1 were found to be low in murine cell lines (~5% cells were positive) whereas in human cell lines basal levels were higher (~15% for MDA-MB-468 and ~25% for Hs578T). However, post-virus infection all cell lines demonstrated an increase in the PD-L1 expression (Figure 1(a, b)). Furthermore, to determine if this holds true for in vivo condition, mice bearing E0771 orthotopic tumors were injected intra-tumorally with PBS or CF33-hNIS-ΔF14.5 and 7 days later tumor sections were stained for PD-L1. Higher levels of PD-L1 were observed in the virus-treated tumors compared to PBS-treated tumors (Figure 1(c)). However, due to higher variation within the groups, the difference in PD-L1 levels did not reach statistical significance (Figure 1(d)).

We also compared virus growth and resulting cytotoxicity in human and murine TNBCs. Not surprisingly, murine TNBCs were found to be less susceptible to the virus compared to the human TNBCs. Murine TNBCs supported little to no growth of the virus whereas human TNBCs highly supported the growth of the virus with approximately 4 logs increase in virus titers 72 h post-infection (Figure 2(a)). Commensurate with the virus growth, human TNBCs were more susceptible to virus-mediated killing compared to the murine TNBCs (Figure 2(b)). We have previously reported similar finding showing discrepancies in growth and cytotoxic ability of CF33 in other human and murine cancer cell lines.\(^{21}\)

**CF33-hNIS-ΔF14.5 in combination with αPD-L1 shows synergistic anti-tumor effect when both agents are injected intra-tumorally**

Since all the tested TNBCs showed up-regulation of PD-L1 in response to virus infection, we hypothesized that a combination of CF33-hNIS-ΔF14.5 with an inhibitor of the PD-L1 will increase the overall therapeutic efficacy. To test the therapeutic effect of the combination we used E0771 syngeneic tumor model because this model has been well characterized as murine TNBC; it is negative for ER, PR and HER2 and has mutated p53.\(^{29-31}\) Furthermore, although E0771 cells make aggressive tumors, in our experience E0771 tumors provide somewhat longer therapeutic time window compared to the extremely aggressive 4T1 murine TNBC model. E0771 cells were used to generate bilateral orthotopic tumors in C57BL/6 mice and after the tumors became palpable mice were treated with PBS or α-PD-L1 antibody or CF33-hNIS-ΔF14.5 or a combination of CF33-hNIS-ΔF14.5 and αPD-L1. Only one tumor/mouse

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**Figure 1. Breast cancer cells up-regulate PD-L1 in response to infection by CF33-hNIS-ΔF14.5.** (a) Cells were mock-infected or infected with CF33-hNIS-ΔF14.5 at MOI 3. Eighteen hours post-infection, cells were stained with APC-conjugated PD-L1 antibody or an isotype antibody and analyzed by flowcytometry. (b) Cells were infected and PD-L1 levels determined as in (a) and mean of three independent experiments ± SEM has been plotted. $p$ values were calculated using Student’s t-test. (c) C57BL/6 mice (n = 4/group) bearing orthotopic E0771 tumors were intra-tumorally injected with PBS or $10^7$ PFU of virus on days 1, 3 and 5. On day 7, tumors were harvested and stained for PD-L1 as described in materials and methods. (d) Area in the tumor sections that stained positive for PD-L1 was calculated using QuPath software and compared between the two groups (n = 4/group). Data presented as mean ± SEM. $p$ value was calculated using Student’s t-test.
was injected with the virus and the other tumor (smaller tumor) was left un-injected. For this experiment virus was injected intra-tumorally and α-PD-L1 antibody was injected intra-peritoneally on each of the experimental days 1, 3 and 5 as shown in the treatment scheme (Figure 3(a)). For the injected tumors, the combination group significantly delayed tumor growth compared to the PBS control group whereas no significant difference was observed for single treatment groups (virus or α-PD-L1) compared to PBS group (Figure 3(b)). Growth kinetics of the uninjected tumors looked more or less similar for single and combination treatment groups, however, compared to PBS group statistical significance was achieved only for α-PD-L1 treated group (Figure 3(b)). Mice were euthanized based on tumor burden and survival graph was plotted. In terms of survival, only the combination treatment significantly increased survival of mice compared to PBS-treated mice whereas survival benefits for mice treated with single agents did not reach statistical significance (Figure 3(c)).

We also studied the combination in a separate experiment in which both α-PD-L1 and the virus were injected intra-tumorally. Mice bearing orthotopic E0771 tumor (one tumor/mouse) were injected intra-tumorally with α-PD-L1 or CF33-hNIS-ΔF14.5 or both on each of the experimental days 1, 3 and 5 (Figure 4(a)). All PBS-treated mice had to be euthanized within 32 days due to tumor burden (Figure 4(b)).
Similarly, six out of eight α-PD-L1-treated mice had to be euthanized within the same time period, two mice showed delayed tumor growth and survived longer. However, in the CF33-hNIS-ΔF14.5-treated group 2 out of 8 mice showed complete tumor regression. Interestingly, 50% of mice (4 out of 8) treated with the combination of CF33-hNIS-ΔF14.5 and α-PD-L1 showed no response whereas the other 50% mice achieved complete tumor regression (Figure 4(b)). No overt toxicities or weight loss were seen in mice in any of the treatment groups (Figure 4(c)). Average tumor volume for either of the single treatment group was not significantly different from that for the PBS group whereas the combination group had significantly lower tumor volume compared to the PBS group (Figure 4(d)). More importantly, the combination treatment showed the highest increase in the survival of the mice (Figure 4(e)). Taken together, CF33-hNIS-ΔF14.5 and α-PD-L1 showed minimal anti-tumor effect in E0771 model when used as a single agent; however, their combination resulted in significant anti-tumor effect and prolonged survival of the treated mice, and the anti-tumor efficacy was more pronounced when both treatment agents were administered intra-tumorally.

**Immune modulation by CF33-hNIS-ΔF14.5 in TME**

Tumors treated intra-tumorally with CF33-hNIS-ΔF14.5 alone or in combination with αPD-L1 antibody were found to have higher levels of immune cells (CD45+) compared to PBS-treated tumors on day 7 after the first dose of treatment (Figure 5(b)). Massive infiltration of CD8+ T cells was observed in the tumors treated with CF33-hNIS-ΔF14.5 alone or in combination with αPD-L1 antibody (Figure 5(b) & Figure 6). However, there was no significant difference in the levels of CD8+ T cells in PBS and αPD-L1-treated mice. We also analyzed CD8+ T cells for their activation status using IFNγ staining. CF33-hNIS-ΔF14.5-treated tumors showed a trend of higher IFNγ+ CD8+ T cells however this increase did not reach statistical significance compared to PBS-treated tumors. Among all the treatment groups only the combination treatment showed increased immune cells and IFNγ expression.
resulted in significantly higher numbers of IFNγ+ CD8+ T cells compared to the PBS group. While there was a trend of higher CD4+ T cells in the combination treatment group compared to that in the PBS group, the difference was not significant due to high variation within the PBS group (Figure 5(b)). Next, we also performed IHC staining on tumor sections and quantified CD4+ and CD8+ T cells to verify the results from FACS analysis. Whole tumor
sections were scanned, and quantification was performed using QuPath software. Results from IHCs were found to be similar to the results from FACS analysis (Figure 6).

**Combination treatment elevates pro-inflammatory cytokines in TME but not in serum**

In TME, cytokines are secreted mainly by immune cells and stromal cells.

Cytokines play important roles in the proliferation, differentiation, activation and migration of immune cells. In order to determine the effect of treatments on cytokine in the TME, we measured the levels of different cytokines in tumor lysates 7 days after the first dose of treatment. We used a kit that could simultaneously measure 13 cytokines (IFNγ, IL-5, TNFα, IL-2, IL-6, IL-9, IL-17A, I-17F, IL-21, IL-22 and IL-13), which are collectively secreted by Th1, Th2, Th9, Th17, Th22 and T follicular cells. Out of the 13 cytokines analyzed, only two cytokines IL-6 and IFNγ were found to be significantly different among the treatment groups (Figure 7(a,c)). Compared to the PBS group, levels of IL-6 were found to be significantly higher only in the combination group but the levels of IFNγ were found to be significantly higher both in the CF33-hNIS-ΔF14.5 treatment group and the combination treatment group. We also measured the levels of those 13 cytokines in the serum of the treated mice (Figure 7(b)). Interestingly, there was no significant difference in the levels of any cytokines among the treatment groups suggesting that the effect of treatments on cytokine levels was mostly confined to the TME. Levels of the pro-inflammatory cytokines IFNγ, IL-6 and TNF-a were much lower in serum compared to those in the TME (Figure 7(c)).

**CF33-hNIS-Δ14.5 alone or in combination with αPD-L1 induces anti-tumor immunity in treated mice**

Immunotherapy often results in the induction of anti-tumor immunity in responding subjects thus providing long-term protection against the malignancy. In pre-clinical models, ‘tumor re-challenge’ is considered the gold standard test for the development of anti-tumor immunity, in which animals with complete tumor-regression are re-injected with the same tumor cells. Existence of immunity against those tumor cells would prevent tumor development whereas animals lacking specific immunity against the tumor cells would develop tumors. We performed ‘tumor re-challenge’ experiments to

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**Figure 7.** CF33-hNIS-Δ14.5 in combination with αPD-L1 increases pro-inflammatory cytokines in TME but not in serum. E0771 tumors were treated as in Figure 4 and tumors as well as serum were harvested 7 days after treatment. Tumors were homogenized to obtain tumor lysates. A multiplex bead-based assay was used to quantify 13 cytokines in the tumor lysates. (a) Heat-map showing all the 13 cytokines in the tumor lysates in the 4 treatment groups (n = 4 mice/group). (b) Heat-map showing all the 13 cytokines in the serum of the mice. Samples were analyzed in duplicates. (c) Mean concentration of IFNγ, IL-6 and TNF-a in tumor lysates and serum have been plotted with SEM. One-Way ANOVA was used to calculate p values. Note: color scales for Tumor (Figure (a)) and Serum (Figure (b)) are different.
determine if the ‘cured’ mice had developed anti-tumor immunity. Age-matched naïve mice were used as control. Mice were injected with E0771 cells in the mammary fat pad (on opposite side) and were monitored for tumor growth. All naïve mice had palpable tumors 2 weeks after injection of cells and by 6 weeks all those mice had to be euthanized due to tumor burden (Figure 8(a,b)). However, mice that were ‘cured’ of their tumors did not develop tumors upon re-challenge and continued to survive. To confirm E0771 tumor-specific immunity, splenocytes collected at the end of experiments were re-stimulated with E0771 cells and levels of CD107+ CD8 T cells were compared among naïve mice and the mice that were ‘cured’ of their tumors. Higher levels of CD107+ CD8 T cells were found in the ‘cured’ mice compared to that in the naïve mice (Figure 8(c)). Together, these data suggest that the mice showing complete response to the therapy developed tumor-specific immunity.

Discussion

Oncolytic viruses have shown promising results in multiple malignancies in preclinical and clinical studies. In addition to other modes of action, OVs stimulate anti-tumor immunity.9,39–42 There are several mechanisms through which OVs could stimulate the immune system. First, oncolysis by OVs may release tumor-specific antigens leading to the activation of the adaptive immune system.43–46 Second, OVs could induce immunogenic death of cancer cells, which results in the activation of immune system against tumor cells.22,23 We have previously reported that CF33 induces immunogenic cell death in multiple tumor cell lines.21,26 In this study, we found that breast cancer cells, both human and murine, upregulate PD-L1 expression in response to infection by CF33-hNIS-ΔF14.5. This increase in the PD-L1 levels could be the consequence of virus-induced type-1 interferons. Our data is in congruence with a study published by Liu et al. which showed upregulation of PD-L1 in a panel of human and murine cell lines encompassing a variety of malignancies after infection with an oncolytic vaccinia virus.27 The study by Liu et al. showed that the combination of an oncolytic vaccinia virus with αPD-L1 antibody results in synergistic anti-tumor effect in mouse models of colon

Figure 8. Mice treated with CF33-hNIS-ΔF14.5 alone or in combination develop tumor-specific immunity. Mice that underwent complete tumor regression from Figure 4, and remained tumor free for 30 days, were re-challenged with 10^3 E0771 cells in the abdominal mammary fat pad of opposite side. Age-matched naïve mice were used as control. (a) Tumor volumes were measured weekly and average tumor volumes have been plotted with SEM. Tumors were detected only in the naïve mice. (b) Mice were euthanized when their tumor exceeded 2500 mm^3 in volume, and survival was compared using Kaplan–Meier survival plot. (c) CD8+ T cells were isolated from the mice at the end point and re-stimulated with E0771 cells. Cells were stained with antibodies against CD8 and CD107 and analyzed using flow cytometry. Per cent CD8+ T cells positive for CD107 marker have been plotted with SEM and compared among the groups. Stats: One-Way ANOVA.
that model in this study. Nevertheless, CF33-hNIS-ΔF14.5 was found to favorably modulate immune cells and cytokines in the TME of E0771 syngeneic model and showed some anti-tumor efficacy with complete tumor regression in 2 out of 8 mice. CF33-hNIS-ΔF14.5 significantly increased tumor infiltration by CD8+ T cells and there was a trend of increased PD-L1 expression on nonimmune cells (CD45-) in virus-treated tumors. The checkpoint inhibitor aPD-L1 has been shown to work better in tumors with high levels of CD8+ T cells and high levels of PD-L1 on tumor cells.53,55,56 Hence, we combined CF33-hNIS-ΔF14.5 with aPD-L1 antibody and the combination resulted in much better anti-tumor efficacy compared to single treatment with either components. In this study, we injected virus intra-tumorally and the aPD-L1 antibody either intra-tumorally or intra-peritoneally. The reason for the intra-tumoral injection of the aPD-L1 antibody was to study the practicability of modifying our virus CF33-hNIS-ΔF14.5 to encode aPD-L1 antibody. Furthermore, since OVs could potentially evoke autoimmune responses, systemically delivered checkpoint inhibitors may result in higher levels of autoimmune side-effects.41,57,58 Therefore, in context of the combination of OV and checkpoint inhibitors considering intra-tumoral injection of checkpoint inhibitor may not be unreasonable. Our data show that intra-tumoral injection of the virus alone or in combination with aPD-L1 antibody increases pro-inflammatory cytokines IFNγ and IL-6 in TME but not in the serum. The fact that pro-inflammatory cytokines were increased only in tumors, and their levels were much lower in serum, suggests that the treatment specifically induces immunologically conducive or ‘hot’ environment within the tumors. Interestingly, in our study the combination of CF33-hNIS-ΔF14.5 virus and aPD-L1 antibody worked better when both agents were delivered intra-tumorally compared to the combination where the virus was delivered intra-tumorally and the antibody was delivered intra-peritoneally. While the former combination resulted in complete tumor regression in 50% of mice, no complete tumor regression was achieved in any mouse in the later combination treatment. Also, in the later combination group using bilateral tumor model we found that while the virus-injected tumors showed significantly slower growth compared to the PBS group, the un-injected tumors despite slower growth did not reach statistical significance. Of note, in this study we have tested only one dose of virus (10⁷ PFUs) and one dose of aPD-L1 antibody (100 μg), both of which are lower doses compared to doses used in previous studies. For example, most published studies have used 200 μg or higher doses of aPD-L1 antibody27,59-61 and/or 10⁸ PFUs of oncolytic vaccinia virus in immune-competent mice.25,62,63 Therefore, it is likely that better therapeutic efficacy could be achieved in the studied model if higher doses of the therapeutic agents are administered. Nevertheless, mice that underwent complete tumor regression after intra-tumoral treatment with CF33-hNIS-ΔF14.5 alone or the combination of CF33-hNIS-ΔF14.5 and aPD-L1 were protected from tumor development when re-challenged with E0771 cells, suggesting that those animals developed anti-tumor immunity against the tumor cells as a result of treatment. This finding was further confirmed by CD107 expression by splenic CD8+ T cells after restimulation with E0771 cells.

In summary, our data show that CF33-hNIS-ΔF14.5 increases PD-L1 levels on breast cancer cells and favorably modulates the immune landscape in E0771 tumors setting the stage for the activity of the immune checkpoint inhibitor aPD-L1. Hence, the combination of CF33-hNIS-ΔF14.5 with aPD-L1 showed better therapeutic efficacy in the studied tumor model. Addition of other checkpoint inhibitors such as anti-CTLA-4 may further enhance the overall anti-tumor efficacy.

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