Improved Antitumoral Efficacy of Mesothelin Targeted Immune Activating Fusion Protein in Murine Model of Ovarian Cancer

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Abstract
The candidate therapeutic fusion protein (scFv-MtbHsp70) is a recombinant Mycobacterium tuberculosis heat shock protein 70 (MtbHsp70) fused with a single chain antibody (scFv) targeting mesothelin, combining the immune-targeting capacity of the scFv with the broader immune activating capabilities of MtbHsp70. The previous version of the fusion protein, VIC-007, markedly enhanced survival of ovarian tumor-bearing mice through the augmentation of tumor-specific cell-mediated immune responses. In this study, a new version of the fusion protein, VIC-008, was reconstructed from VIC-007 by slight modifications that removed redundant amino acids and introduced a single amino acid mutation, phenylalanine to valine, at position 381 of MtbHsp70 to prevent non-specific peptide binding and presentation while retaining immune-stimulatory capacity. VIC-008 showed significantly improved protection in control of tumor growth and prolongation of the survival of tumor-bearing mice compared to VIC-007. These findings offer a definitive preclinical validation of VIC-008 as a therapeutic agent for ovarian cancer.

Keywords
Mycobacterial Hsp70, Mesothelin, Single Chain Variable Fragment, Ovarian Cancer

Introduction
Ovarian cancer is the most lethal malignancy in women, with 22,280 new cases and 15,460 deaths estimated in the United States in 2012 [1]. Surgical debulking followed by platinum-based chemotherapy remains the current standard-of-care treatment to which approximately 80% of patients respond favorably. However, more than 60% of patients who initially achieve remission eventually relapse and less than 40% survive beyond 5 years [2]. Thus, novel treatments for ovarian cancer are urgently needed. Although clinical trials of various immunotherapeutic modalities have not yet yielded significant benefit [3], the failures to date can be understood in light of the complexities of immune regulation, potentially leading to the development of more efficacious immunotherapies.

A novel fusion protein, VIC-007, consists of the broadly immune-activating Mycobacterium tuberculosis-derived heat shock protein 70 (MtbHsp70) and the tumor antigen targeting activity of a single-chain variable fragment (scFv) binding mesothelin (MSLN), validated in a syngeneic, orthotopic and immune competent murine model of ovarian cancer [4-6]. The fusion mechanism of anti-MSLN scFv and MtbHsp70 takes advantage of the immune-activating action of MtbHsp70 and the tumor-targeting activity of the scFv, which will yield anti-tumor responses against the broadest profile of tumor antigens.

Although previous studies showed that VIC-007 significantly enhanced survival of immune competent mice with ovarian or malignant mesothelioma tumors through the augmentation of tumor-specific cell-mediated immune responses [14], the fusion protein did not result in long-term remission. In this study a new version of the fusion protein, VIC-008, was reconstructed from VIC-007 to remove redundant amino acids and minimize the activity of the natural peptide-binding site of MtbHsp70. We compared VIC-007 and VIC-008 side by side in the same set of mice and found that VIC-008 conferred significantly improved antitumoral efficacy in a syngeneic, orthotopic and immune competent murine model of ovarian cancer.
Vic-007

GSS

RGLTRYTSRRNYTDNATVYVSRSKSMIPDSRKEQRLGNSVQFEDTVXTVC

AMGKMTXYYSMQVWQNTTTSQVDYTVSGILSOGSVSSGSGSSGSSG

SLSASFSGASLCTTCLSGPRVYWYWRQQGPFSQNYLNNSEDSCK

QQSGSVPFSFSGKDAANAYQVPGISLGRRSEHEDAYCMWHSSAAYFGG

TQLTVL$

SGNLQEGPOQGSSGSGSGSGSGASAGHNRGAAAAYUGIDGTVNYV

SWYSGVVFQVAFFAKNSLVRGQFQAVAVFVNGYRTVR

SVKRYHKSMSWSWIDVISKKYQTEPSIARILMLKRDAYALEIDTVAITV

PANFQDNAATRDAQGIQNLMVRNENEPAAYLAAYLGDKGEKEQRIV

LVDGQQVRIEFLTIFEESPSTTTDAQDNPQVYQQEGERAIAH

GDDDQSFQNMDTSGVHFMDVAVQAALQVLQSVGVLVDVFTLSL

CITVKEQVRIEFLTIFESPSTTTDAQDNPQVYQQEGERAIAH

VNKLQSGFETGLTPAPFIQPOPQVFSTDIDAVIHGTQDKTGMREIKI

QKEDGSLAEDIDMRKLAAHAAEIREEKREDVQRNVLTQVFYKE

QREAMGGKVEPDVNLVCQAADDVVKAEKLQGSGS

CQAQVYAAQAASATQAAC#HPGEFGEPHASGADDVDVEAVDDGREAK

Vic-008

VQTVQQGPSGGPVTFQSTLLICAISSGDSWSSNSATWQIGPGSRELELMG

RTYRSRKYWDNVAYDSRSFQMDPPKVNLFNITVQAYT GMTYVMDQNQGFTVTSQGSLGSGGSQSGSSLQPVSQSSL$SASASGASLSLCTTCLSGPRVYWYWRQQGPFSQNYLNNSEDSCK

SOFSPRSFGKDSADANAGVLLILLTRSEADAVCMWHSSAAYFGG

TVLGGQVVFQVAFFAKNSLVRGQFQAVAVFVNGYRTVR

SVKRYHKSMSWSWIDVISKKYQTEPSIARILMLKRDAYALEIDTVAITV

PANFQDNAATRDAQGIQNLMVRNENEPAAYLAAYLGDKGEKEQRIV

LVDGQQVRIEFLTIFEESPSTTTDAQDNPQVYQQEGERAIAH

GDDDQSFQNMDTSGVHFMDVAVQAALQVLQSVGVLVDVFTLSL

CITVKEQVRIEFLTIFESPSTTTDAQDNPQVYQQEGERAIAH

VNKLQSGFETGLTPAPFIQPOPQVFSTDIDAVIHGTQDKTGMREIKI

QKEDGSLAEDIDMRKLAAHAAEIREEKREDVQRNVLTQVFYKE

QREAMGGKVEPDVNLVCQAADDVVKAEKLQGSGS

CQAQVYAAQAASATQAAC#HPGEFGEPHASGADDVDVEAVDDGREAK


displayed by IVIS Spectrum.

imaged by IVIS Spectrum.

kg body weight of D-luciferin 10 min in advance and subsequently

live imaging by IVIS Spectrum

tumor cell inoculation using

streptomycin, and 10% fetal bovine serum in humidified atmosphere

DMEM with 2 mmol/L L-glutamine, 10 units/ml penicillin, 10 μg/ml

luciferase, here named Luc-ID8. Cells were maintained at 37°C

transfected with luciferase lentiviral vector and stably expressed

(University of Kansas Medical Center, Kansas City, KS) [15], were

protocols that were approved by the Massachusetts General Hospital

were performed in a manner that was blinded to the observer under

was followed by 3 further treatments at 7-day intervals. All studies

days after tumor cell inoculation with i.p. injections of VIC-007 (4

Jackson laboratories. Mice with ovarian tumors were treated 7

weeks post-inoculation with tumor cell inoculation


treated with Luciferase using for animal injections.

with Trypsin EDTA (Mediatech) for animal injections.

In vivo imaging of tumor growth

Intraperitoneal tumor growth was monitored weekly after

tumor cell inoculation using in vivo live imaging by IVIS Spectrum

(ParkinElmer). Mice were injected intraperitoneally with 150 mg/

kg body weight of D-Luciferin 10 min in advance and subsequently

imaged by IVIS Spectrum.

Mouse survival

For survival studies, we observed the mice daily 1 week after

inoculation of tumor cells. Tumor generations were consistently first
evident via abdominal distension secondary to malignant ascites, and tumor-bearing mice were euthanized at the endpoint when there

were signs of distress, including fur ruffling, rapid respiratory rate, hunched posture, reduced activity, and progressive ascites formation as previously described [16].

**Statistical analysis**

Statistical differences between three or more experimental groups were analyzed using Two-Way ANOVA, followed by Tukey’s

multiple comparison test when mean of each group is compared with that of every other group. Survival was analyzed with the Log-

rank test. Prism 6.0 software (GraphPad Software) was used for all the statistical analysis.

**Results and Discussion**

Reconstruction of the fusion protein scFv-MtbHsp70

The fusion protein scFv-MtbHsp70 was constructed with Vh and

Vl from anti-MSLN p4 scFv [17] fused to full length MtbHsp70 with

a (GAS)3 linker in between, which has been shown in our previous study [14]. The previous version of the fusion protein VIC-007

achieved significant control of tumor growth and prolongation of the survival of tumor-bearing mice, but the antitumoral efficiency of the treatment regimen used needed to be improved. Antigenic peptides linked to MtbHsp70 through both non-covalent binding and by genetic fusion can elicit both MHC class I-restricted CD8+ and MHC class II-restricted CD4+ T-cell responses [18-22]. In this study we

developed a new version of the scFv-MtbHsp70 fusion protein, fusion protein, VIC-008, which was modified from the original VIC-007 by the elimination of redundant amino acids and the introduction of a single amino acid mutation, phenylalanine (F) in place of valine (V), at position 381 of MtbHsp70 (Figure 1). This change is designed to prevent peptide binding [23] while retaining the immune-stimulatory capacity of the protein, in order to reduce the possibility that MtbHsp70 might incidentally bind and deliver other antigens that could result in off target effects or the induction of tolerance or autoimmunity.

The fusion proteins were constructed and expressed by WuXi App Tech (Shanghai, China) in CHO cells and provided at a purity of above 95% by HPLC and an endotoxin level of less than 1.0 EU/mg.

**VIC-008 enhances the control of tumor growth**

Murine ovarian cancer was established by i.p. injection of

syngeneic cancer cells Luc-ID8 in immune competent C57BL/6 mice and treated with VIC-007 and VIC-008 as described in the section of materials and methods. As shown in figure 2, both VIC-007 and VIC-008 significantly slowed tumor growth as recorded by bioluminescence signals compared to saline (p < 0.0001 and p < 0.0001) while VIC-008 further significantly delayed tumor growth compared to VIC-007 (p < 0.0001).

**VIC-008 enhances the prolongation of mouse survival**

We further evaluated the efficacy of VIC-007 and VIC-008 to

prolong survival in the tumor-bearing mice. As shown in figure 3, both VIC-007 and VIC-008 significantly enhanced the survival of

tumor-bearing mice compared to saline (p = 0.0253 and p = 0.0002)

with increased median survival of 55 days from saline to 60 days from VIC-007 and further to 65 days from VIC-008. VIC-008 further significantly prolonged the survival of the tumor-bearing mice compared to VIC-007 (p = 0.0301).

Taken together, these data showed that the new version of the fusion protein VIC-008 significantly delayed the tumor growth and

prolonged the survival in a syngeneic murine model of ovarian cancer. Improved mouse survival of VIC-008 compared to VIC-007 is likely related to the changes made to the protein sequences. Since the efficacy of immunotherapeutic agents is often limited by negative immunological regulations that attenuate immune responses in the tumor microenvironment when used as a single agent, we will conduct further testing of VIC-008 in combination with checkpoint blockade.

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**Materials and Methods**

**Cells**

The ID8 ovarian cancer cells, a kind gift from Kathy Roby (University of Kansas Medical Center, Kansas City, KS) [15], were transfected with luciferase lentiviral vector and stably expressed luciferase, here named Luc-ID8. Cells were maintained at 37°C in DMEM with 2 mmol/L L-glutamine, 10 units/ml penicillin, 10 μg/ml streptomycin, and 10% fetal bovine serum in humidified atmosphere with 5% CO2. Cells were cultured until 80% confluent, and harvested with Trypsin EDTA (Mediatech) for animal injections.

**Animal model and treatment**

Ovarian cancer was established by Intraperitoneal (i.p.) injection of syngeneic cancer cells Luc-ID8 (5 × 105 cells per mouse) into 6-week-old female C57BL/6 mice. All mice were purchased from Jackson laboratories. Mice with ovarian tumors were treated 7 days after tumor cell inoculation with i.p. injections of VIC-007 (4 μg per mouse), VIC-008 (4 μg per mouse), or normal saline. This was followed by 3 further treatments at 7-day intervals. All studies were performed in a manner that was blinded to the observer under protocols that were approved by the Massachusetts General Hospital Subcommittee on Research Animal Care (SARC).

**In vivo imaging of tumor growth**

Intraperitoneal tumor growth was monitored weekly after tumor cell inoculation using in vivo live imaging by IVIS Spectrum (PerkinElmer). Mice were injected intraperitoneally with 150 mg/kg body weight of D-Luciferin 10 min in advance and subsequently imaged by IVIS Spectrum.

**Mouse survival**

For survival studies, we observed the mice daily 1 week after
that are capable of reprogramming suppressive and stimulatory signals in the intratumoral environment that together with VIC-008 has the potential to yield more powerful cancer control.

Conclusion

Our study provides a definitive preclinical validation of the mesothelin targeted immune activating fusion protein as a therapeutic agent for ovarian cancer and supports the continued exploration of this novel fusion protein alone or in combination with immune checkpoint inhibitors or chemotherapy for MSLN-expressing cancers.

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Competing Interests Statement

We declare that we have no competing financial interests.

Abbreviations

MSLN: mesothelin; scFv: single-chain antibody variable fragment; MTB: Mycobacterium tuberculosis; Hsp: heat shock protein; DC: dendritic cell; i.p.: intraperitoneal.

References


