A Virus-Like-Particle immunotherapy targeting Epitope-Specific anti-xCT expressed on cancer stem cell inhibits the progression of metastatic cancer in vivo

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ABSTRACT
Aggressive forms of breast cancer, such as Her2
+ and triple negative breast cancer (TNBC), are enriched in breast cancer stem cells (BCSC) and have limited therapeutic options. BCSC represent a key cellular reservoir for relapse, metastatic progression and therapeutic resistance. Their ability to resist common cytotoxic therapies relies on different mechanisms, including improved detoxification. The cystine-glutamate antiporter protein xCT (SLC7A11) regulates cystine intake, conversion to cysteine and subsequent glutathione synthesis, protecting cells against oxidative and chemical insults. Our previous work showed that xCT is highly expressed in tumorspheres derived from breast cancer cell lines and downregulation of xCT altered BCSC function in vitro and inhibited pulmonary metastases in vivo. We further strengthened these observations by developing a virus-like-particle (VLP; AX09-0M6) immunotherapy targeting the xCT protein. AX09-0M6 elicited a strong antibody response against xCT including high levels of IgG2a antibody. IgG isolated from AX09-0M6 treated mice bound to tumorspheres, inhibited xCT function as assessed by reactive oxygen species generation and decreased BCSC growth and self-renewal. To assess if AX09-0M6 impacts BCSC in vivo seeding, Her2
+ TUBO-derived tumorspheres were injected into the tail vein of AX09-0M6 or control treated female BALB/c mice. AX09-0M6 significantly inhibited formation of pulmonary nodules. To evaluate its ability to impact metastases, AX09-0M6 was administered to mice with established subcutaneous 4T1 tumors. AX09-0M6 administration significantly hampered tumor growth and development of pulmonary metastases. These data show that a VLP-based immunization approach inhibits xCT activity, impacts BCSC biology and significantly reduces metastatic progression in preclinical models.

Introduction

It is estimated that in 2017, 252,000 women will be diagnosed with breast cancer in the United States including 30,000 new cases of triple negative breast cancer (TNBC), a highly aggressive form of breast cancer.1 According to the hierarchical cancer model, mammary tumors, as the majority of cancers, are organized in a cellular hierarchy with cancer stem cells (CSC) at the apex.2 CSC are often enriched in highly metastatic breast cancers, including TNBC and Her2
+ breast cancer.3 CSC have the unique biological properties necessary for maintenance and spreading of the tumor, and through asymmetric division can differentiate into cancer cells that compose the tumor bulk.4,5 Due to their resistance to traditional radiation and chemotherapies,6 CSC represent a reservoir for the relapse, metastatic evolution and progression of the disease after treatment. Therefore, successful eradication of CSC represents a major barrier towards effective cancer treatments.

The ability of CSC to resist common cytotoxic therapies relies on different mechanisms, including improved detoxification ability. The cystine-glutamate antiporter protein xCT (SLC7A11), the light chain of the antiporter system xCT
−, regulates cystine intake, conversion to cysteine and subsequent glutathione (GSH) synthesis, protecting cells against oxidative and chemical insults.7,8 xCT is a 12 multi-pass transmembrane protein, characterized by 6 extracellular domains (named ECD1-6), in which the N- and C-termini are located intracellularly.9 xCT expression is low in normal cells and highly restricted to a few cell types (astrocytes, microglia and other subsets of myeloid cells),10,11 but elevated levels of xCT protein are observed in a high percentage of mammary tumors including TNBC.12 High levels of xCT mRNA and protein correlate with significant reduction in distal metastases-free and overall survival.13,14 xCT expression is upregulated in breast CSC (BCSC)15 and other solid tumor stem cells,16-21 and several studies show that xCT physically interacts with the
well-known stem cell proteins, CD44 and MUC1. The frequency of xCT expression on a variety of CSC suggests that therapies targeting xCT may be effective in treating many tumors with high stem cell frequencies including gastrointestinal and pancreatic cancers. Our published work established xCT as an immunotherapeutic target for BCSC. Microarray analysis identified xCT as a gene that was upregulated in BCSC and subsequent studies confirmed xCT upregulation in human and murine BCSC. Inhibition of xCT function resulted in a significant decrease of BCSC viability, reduced self-renewal and altered cellular redox levels. A direct role for xCT in breast cancer growth and metastasis was shown by immunizing mice with a DNA-based vaccine expressing the full-length xCT protein. Administration of the xCT vaccine prior to or in mice with existing tumors resulted in significantly decreased pulmonary metastases and tumors containing significantly less BCSC compared to control animals. The combination of xCT vaccination and chemotherapy synergistically reduced metastatic disease compared to mice that were treated with either therapy alone. Similar results were observed by inhibiting xCT function with the small molecule sulfasalazine by inhibiting xCT function with the small molecule sulfasalazine (SASP), a Food and Drug Administration (FDA)-approved anti-inflammatory drug characterized by low specificity for xCT, short bioavailability and numerous side effects. The mechanism of action for DNA vaccination usually acts through cytotoxic CD8 T cell activation. However, there was no T-cell response to the major xCT T-cell epitope in DNA vaccinated mice, and immunization in B-cell deficient mice showed inadequate therapeutic responses. IgG antibodies isolated from the sera of vaccinated mice inhibited BCSC function in vitro in a similar manner as siRNA and SASP treatment, demonstrating that vaccine-induced xCT antibodies represented the therapeutic effectors. However, antibody titers achieved using the DNA vaccine were low, suggesting that the development of new therapies generating a focused, high titer antibody response would lead to greater inhibition of metastatic progression.

The virus-like particle (VLP) vaccines generate strong immune responses due to their optimal size, particulate nature, and potent intrinsic adjuvant activity. Agilvax’s VLP technology is derived from a family of single-stranded RNA bacteriophages, including MS2, PP7, Qβ, and AP205. To create a VLP that was more thermodynamically stable and dramatically more tolerant of foreign AB loop peptide insertions, Agilvax engineered a single-chain dimer version of the MS2 coat protein. Resulting VLPs are comprised of a single coat protein that self-assembles into a 27 nm diameter icosahedral particle consisting of 90 coat protein dimers. Our VLP technology allows for the precise control of epitope size, structure (loop/linear) and valency to optimize immune responses for a specific antigen. Epitopes are displayed in an ordered, geometric pattern on the surface of the VLPs and elicit robust antibody responses, even against self-antigens. In this paper, we produced and tested a novel VLP-based immunotherapy (AX09-0M6) that displays the 6th extracellular domain (ECD6) of human xCT structure to display on the surface of our VLP technology. The various ECD of human xCT were cloned into the coat protein (CP) AB loop rather than in a linear display point of our VLP to more closely mimic the native cellular structure of the xCT domains. The VLPs were expressed in E. coli, purified and initial characterization showed that VLPs displaying ECD1, ECD3 and ECD6 assembled properly. Based upon ECD6 full homology in human and mouse xCT sequence, the VLP displaying the ECD6 peptide (AX09-0M6) was used for the pre-clinical studies reported here.

Results

Construction of xCT VLPs

To construct a VLP-based immunotherapy targeting xCT, we identified five (out of six) potential ECD from the human xCT structure to display on the surface of our VLP technology. Our published work established xCT as a potential target for BCSC. Microarray analysis identified xCT as a gene that was upregulated in BCSC and subsequent studies confirmed xCT upregulation in human and murine BCSC. Inhibition of xCT function resulted in a significant decrease of BCSC viability, reduced self-renewal and altered cellular redox levels. A direct role for xCT in breast cancer growth and metastasis was shown by immunizing mice with a DNA-based vaccine expressing the full-length xCT protein. Administration of the xCT vaccine prior to or in mice with existing tumors resulted in significantly decreased pulmonary metastases and tumors containing significantly less BCSC compared to control animals. The combination of xCT vaccination and chemotherapy synergistically reduced metastatic disease compared to mice that were treated with either therapy alone. Similar results were observed by inhibiting xCT function with the small molecule sulfasalazine (SASP), a Food and Drug Administration (FDA)-approved anti-inflammatory drug characterized by low specificity for xCT, short bioavailability and numerous side effects. The mechanism of action for DNA vaccination usually acts through cytotoxic CD8 T cell activation. However, there was no T-cell response to the major xCT T-cell epitope in DNA vaccinated mice, and immunization in B-cell deficient mice showed inadequate therapeutic responses. IgG antibodies isolated from the sera of vaccinated mice inhibited BCSC function in vitro in a similar manner as siRNA and SASP treatment, demonstrating that vaccine-induced xCT antibodies represented the therapeutic effectors. However, antibody titers achieved using the DNA vaccine were low, suggesting that the development of new therapies generating a focused, high titer antibody response would lead to greater inhibition of metastatic progression.

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AX09-0M6 administration induces a functional xCT antibody response

To evaluate antibody responses, female BALB/c mice were immunized two times in the absence of exogenous adjuvant.
with AX09-0M6 or control MS2 wild-type (MS2 wt) and a peptide ELISA was used to measure the end-point titer. Sera from AX09-0M6 treated BALB/c mice were serially diluted and incubated with xCT ECD6 peptide coated wells and specific antibody binding was analyzed by ELISA. AX09-0M6 generated peptide antibody titers from $5 \times 10^2$ to $1.4 \times 10^4$ with an average of $3 \times 10^3$ (Fig. 2A), whereas no detectable binding was observed in mice treated with MS2 wt immune sera. AX09-0M6 was also analyzed for relative amounts of different IgG isotypes. AX09-0M6 induced high levels of IgG2a compared to IgG1, indicative of a Th1 biased antibody response (Fig. 2B). IgG2b levels were also elevated compared to IgG1 (Fig. 2B), while little detectable IgG3 was observed (data not shown).

To analyze the ability of AX09-0M6 induced antibodies to bind BCSC, tumorspheres derived from TUBO (mouse HER2$^+$; Fig. 2C), 4T1 (mouse TNBC; Fig. 2D), HCC-1806 (human TNBC; Fig. 2E) and MDA-MB-231 (human TNBC; Fig. 2F) cells were incubated with IgG purified from sera of mice vaccinated with AX09-0M6 or MS2 wt and analyzed by FACS. A commercial anti-xCT Ab was used as a control. IgG from AX09-0M6 vaccinated mice bound to both mouse and human xCT$^+$ BCSC, while no signal was observed with IgG from MS2 wt treated mice.

These results were confirmed by the ability of IgG purified from AX09-0M6-immunized mouse sera to stain tumorspheres derived from TUBO (Fig. 3A), 4T1 (Fig. 3B), HCC-1806 (Fig. 3C) and MDA-MB-231 (Fig. 3D) cells, as shown by confocal images (Fig. 3, on the left) and fluorescent signal quantification graphs, calculated as integrated density normalized to nuclei number (Fig. 3, on the right). No binding was observed in cells incubated with purified IgG from MS2 wt treated mice.

**Antibodies induced by AX09-0M6 inhibit BCSC function**

To test the effect exerted by anti-xCT antibodies induced by VLP vaccination on BCSC, tumorspheres generated from TUBO cells were incubated for 5 days with IgG purified from the sera of vaccinated mice or with the xCT inhibitor sulfasalazine (SASP). As reported in Fig. 4A, IgG from MS2 wt treated mice did not affect BCSC, while those purified from AX09-0M6 vaccinated mice reduced the number (Fig. 4A, panel i) and dimension of spheres (Fig. 4A, panel ii and Supplementary Fig. 1A), exerting an effect comparable to SASP. This reduced sphere-generation ability was accompanied by both a decrease in the percentage of BCSC, evaluated as cells positive for the Aldefluor reagent that measures aldehyde dehydrogenase-1 activity (Fig. 4A, panel iii), and an increase in Reactive Oxygen Species (ROS) content as compared with MS2 wt (Fig. 4A, panel iv).

The ability of anti-xCT antibodies induced by VLP vaccination to inhibit BCSC was not restricted to the TUBO model, as similar results were obtained in other murine (4T1; Fig. 4B and Supplementary Figure 1B) and human (HCC-1806 and MDA-MB-231, Fig. 4C, D and Supplementary Fig. 1C, D) TNBC cell lines. These results suggest that AX09-0M6 elicits anti-xCT antibodies affecting BCSC self-renewal ability.

ROS levels are regulated by xCT via production of GSH, and inhibition of xCT function using a variety of approaches results in an increase in intracellular ROS levels. To investigate if AX09-0M6 elicits antibodies that inhibit xCT function, tumorspheres were incubated for 5 days with IgG purified from MS2 wt or AX09-0M6 vaccinated animals. As seen in Fig. 4 (panel iv), IgG from AX09-0M6 mice resulted in a significant increase in the intracellular ROS levels similar to SASP. In contrast, there were no differences in ROS levels from tumorspheres incubated with IgG from MS2 wt sera, suggesting that AX09-0M6 elicits antibodies that bind to and inhibit xCT function.

**AX09-0M6 treatment has limited or no effect on normal tissues expressing xCT**

Since xCT is expressed at low levels by mouse (but not human) spleen myeloid cells, we analyzed the relative amount of these cell populations in the spleen of vaccinated mice as compared to untreated or MS2 wt treated mice. As shown in Fig. 5A, AX09-0M6 did not induce any reduction in dendritic cells, neutrophils, and macrophages. When xCT$^+$ cells were evaluated, no significant alterations were found (Fig. 5B), but there was a trend towards a reduction in xCT$^+$ neutrophils and macrophages (Fig. 5B). In the case of macrophages, the reduction was primarily found in the M1 and M0 subsets (Fig. 5C).

In the mouse, xCT is also expressed in the central nervous system by astocytes and microglia. In order to exclude brain toxicity in vaccinated mice, we performed a histological and immunohistochemical analysis of brains from AX09-0M6 vaccinated BALB/c mice whose sera endpoint titers are reported in Fig. 2A. Haematoxylin and Eosin (H&E) staining showed no significant morphological alterations and lesions, which suggests that vaccination-induced toxicity was not present. Slight lesions of uncertain meaning, artifacts or agonics, and tiny lymphocytic infiltrates have been detected, but they were distributed in all groups (untreated, MS2 wt and AX09-0M6 treated; data not shown). Immunohistochemical analysis with the F4/80 microglial marker did not reveal any alterations and differences in the AX09-0M6 treated group versus the MS2 wt treated and untreated groups (Fig. 5D-F). Similarly, no significant inflammatory cell infiltrates or any other abnormalities were detected in H&E-stained sections of brains from outbred CD-1 mice (data not shown) in which immunization with AX09-0M6 generated very robust endpoint titers (Supplementary Fig. 2A). The lack of toxicity was also supported by the absence of any detectable behavioural changes, such as increased aggressiveness, narcolepsy or uncoordinated movements, in living AX09-0M6 mice.

**AX09-0M6 attenuates experimental metastases in vivo affecting the immune infiltrate**

To evaluate the ability of AX09-0M6 to inhibit CSC mediated seeding and resuming growth at the metastatic site in vivo, we used a well-characterized preventive mouse cancer model. Female BALB/c mice were vaccinated twice at two-week intervals with MS2 wt or AX09-0M6. One week after final boost, mice were injected intravenously (i.v.) with TUBO-derived tumorspheres and the number of micrometastases was evaluated histologically three weeks later. Mice treated with AX09-0M6 had a significant decrease (~42%) in the number of pulmonary metastases compared to control animals (Fig. 6A).
**Figure 2.** AX09-0M6 induced strong antibody responses. (A) Peptide ELISA of sera from treated BALB/c mice were serially diluted and incubated with xCT ECD6 peptide coated wells. End-point titers were determined by the highest dilution that was 2-fold over background. No titer was observed with control sera. Each dot represents an individual animal treated with AX09-0M6 and results are from 3 independent experiments. (B) AX09-0M6 sera from panel A were analyzed for relative amounts of different IgG isotypes. The relative ratio of each IgG subclass was calculated for each animal by taking the average OD_{450} value from a given IgG subclass (IgGx) divided by the average OD_{450} from total IgG. (C-F) FACS analysis of tumorspheres derived from TUBO (C), 4T1 (D), HCC-1806 (E) or MDA-MB-231 (F) cells incubated with 50 μg/ml IgG purified from sera of MS2 wt or AX09-0M6-vaccinated mice, or with a commercial anti-xCT Ab. Representative histograms are shown. Graphs show the mean ± SEM of the percentage of positive cells set on the base of FITC-conjugated secondary Ab staining and of mean fluorescence intensity from 3 independent experiments. *, P < 0.05; **, P < 0.01; ***, P < 0.001, Student’s t-test.
To see if AX09-0M6 administration could alter the tumor microenvironment, immune infiltrates in lungs were assessed using FACS analysis. As shown in Fig. 6B, AX09-0M6 induced a significant increase in lung infiltrating NK cells and a trend showing an increase in T cells, while no difference in either γδ T cells and NKT cells were observed compared to untreated or MS2 wt vaccinated mice. Of note, both MS2 wt and AX09-0M6 increased the proportion of CD8+ T cells and decreased that of CD4+ T cells when compared to untreated mice (Fig. 6C), suggesting that VLPs may influence the tumor microenvironment independently of the expression of a specific peptide in the CP AB loop. Although not significant, MS2 wt and, to a greater extent, AX09-0M6, decreased the number of Ly6G+Ly6C+ neutrophils and increased both the Ly6G−Ly6C+ monocytes and F4/80+ macrophages compared to untreated animals (Fig. 6D).

The increased number of NK cells in the lungs of AX09-0M6 treated mice and the fact that AX09-0M6 induces xCT

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**Figure 3.** AX09-0M6-induced antibodies target tumorspheres. Representative immunofluorescence images of TUBO (A), 4T1 (B), HCC-1806 (C) and MDA-MB-231 (D)-derived tumorspheres incubated with IgG (50 μg/ml) purified from sera of BALB/c mice vaccinated with MS2 wt or AX09-0M6. The specific signal (green) was detected with an Alexa Fluor488-conjugated anti-mouse secondary antibody. Nuclei were counterstained with DAPI (blue). Magnification 40X, Scale bar, 40 μm **, P < 0.01, Student’s t-test.
targeting antibodies mainly of the IgG2a subclass, were suggestive of a possible role of antibody-dependent cellular cytotoxicity (ADCC) in vaccine-mediated metastatic inhibition. To test this possibility, we performed an ADCC in vitro assay by using xCT⁺ 4T1 cells as targets incubated with the sera (1:50 dilution) from mice treated with AX09-0M6, MS2 wt or left untreated and autologous splenocytes (SPC) as effectors. As shown in Fig. 6E, SPC in the presence of pooled sera from AX09-0M6 vaccinated mice mediated higher ADCC at 200:1 and 100:1 effector/target ratios than sera from MS2 wt treated (P < 0.0595 and P < 0.0468, respectively) or untreated control (P = 0.0297 and P = 0.0033, respectively) mice.

**AX09-0M6 slows in vivo mammary tumor growth and attenuates spontaneous metastases**

To investigate if AX09-0M6 would affect tumor growth and metastatic progression in mice with existing tumors, we used the 4T1 syngeneic therapeutic model. Tumorspheres generated from 4T1 cells were administered subcutaneously (s.c.) into the mammary fat pad. When the tumors were 1–1.5 mm in diameter, mice were vaccinated with AX09-0M6 followed by a boost 2 weeks later. As seen in Fig. 7A, the tumor growth rate (as measured by tumor volume) was significantly slower in the AX09-0M6 treated animals compared to controls. Forty days after tumor challenge, animals were euthanized and the
The number of micrometastases was measured. Similar to the preventative model, treatment with AX09-0M6 significantly inhibited the number of spontaneous pulmonary metastases compared to control animals (Fig. 7B), suggesting that AX09-0M6 might represent a new immunotherapy option for breast cancer patients.

Discussion

Our previous work showed that xCT is upregulated in human and murine BCSC and established xCT as an immunotherapeutic target for BCSC. In this manuscript, we show that an active immunization approach using a VLP-based immunotherapy targeting xCT significantly decreased lung metastases in a therapeutic model of aggressive TNBC breast cancer. Moreover, this vaccination was successful in decreasing HER2BCSC seeding, thus reducing tumor growth at the metastatic site. In the therapeutic setting, AX09-0M6 also decreased the primary tumor growth rate (as assessed by tumor volume) in the xCT positive 4T1 TNBC model, suggesting that AX09-0M6 may have additional activity in the 30% of TNBC patients whose bulk tumor cells are also xCT positive.

The mechanism of action for AX09-0M6 is mostly through the generation of an oligoclonal xCT antibody response and efficacy is therefore independent from MHC expression and can target CSC that are known to downregulate the expression of MHC class I molecules to escape cytotoxic T lymphocyte attack. AX09-0M6 induced antibodies inhibited xCT function in vitro resulting in a significant decrease of BCSC viability, reduced self-renewal and altered cellular redox levels, showing that xCT is required for BCSC function rather than simply representing a biomarker of BCSC. Therefore, AX09-0M6 treatment (and other therapies inhibiting xCT function) should not lead to cancer escape mutants via antigen loss mechanisms.

Besides the direct effect of antibodies on xCT function, it should be noted that AX09-0M6 induced a predominant IgG2a antibody response, a subclass that has strong ADCC and Complement Dependent Cytotoxicity (CDC) activity. Indeed, sera from AX09-0M6 vaccinated mice was able to induce cytotoxicity against xCT+ tumor cells. Moreover, analysis of tumor infiltrating lymphocytes (TILs) showed higher levels of NK cells in the metastatic lungs from AX09-0M6 treated animals, supporting a possible role of ADCC in vaccine-mediated metastatic inhibition.

The increased T cell infiltration in metastatic lungs, even though not statistically significant, is also promising and can be justified by the antibody-mediated inhibition of xCT function. Indeed, it has been shown that xCT inhibition results in a decrease of β-catenin transcriptional activity, which inversely correlates with CD8+T cell infiltration while it is associated with infiltration of T regulatory cells and their survival.

AX09-0M6 is initially being developed as an adjuvant therapy for TNBC, due to the high frequency of xCT expressing BCSC found in these tumors, the high rates of relapse and metastatic progression and the lack of targeted therapies. However, research suggests that AX09-0M6 may be of benefit to patients with other indications. The xCT protein is expressed in BCSC and differentiated breast cancer cells in a subset of patients presenting with a variety of breast cancer subtypes, suggesting that xCT positive patients who become refractive to frontline, targeted breast cancer therapy may benefit from AX09-0M6 treatment. Among the others, pancreatic, gastrointestinal, glioblastoma and colorectal...
CSC express xCT and inhibition of xCT functions in preclinical models of these cancers using SASP showed reduced metastases and tumor growth.

AX09-0M6 has potential as an add-on therapy for conventional and emerging frontline therapies. Many chemotherapeutic agents upregulate xCT protein in more differentiated cancer cells, which conveys resistance to many of these therapies. Therefore, AX09-0M6 inhibition of xCT function in combination, or sequentially, with chemotherapy may elicit clinical benefit by sensitizing bulk tumors to chemotherapeutic agents as well as targeting BCSC. The use of blocking monoclonal antibodies to the checkpoint inhibitor Programmed Death-Ligand 1 molecule (PD-L1) is currently in trials for TNBC. PD-L1 is expressed on human CSC and a preliminary FACS experiment shows that PD-L1 is also expressed on 4T1-derived BCSC (data not shown). The combination of AX09-0M6 therapy with PD-L1 blockade may be more effective in reducing BCSC function while still blocking the PD1-PD-L1 axis in the

Figure 6. AX09-0M6 attenuates metastases in vivo affecting the immune infiltrate and induces antibodies stimulating ADCC. (A) Mice were treated with MS2 wt or AX09-0M6 in the absence of exogenous adjuvant. One week after final administration, P2 tumorspheres derived from TUBO cells were injected into the tail vein of treated mice. 20 days after cell challenge, the lungs were removed, sectioned and the number of micrometastases was determined. Each dot represents a single animal and is the average number of lung metastases from at least 2 sections. Results shown are the aggregate from 3 independent experiments. **, P = 0.0025; ***, P = 0.0025, Mann Whitney test. (B-D) Cytofluorimetric analysis of immune infiltrates in lungs of mice left untreated (white bars) or vaccinated with MS2 wt (black bars) or AX09-0M6 (gray bars). (B) Graph shows the percentage ± SEM of CD45+ cells expressing the markers of T (CD3+CD49b−), NK (CD3−CD49b+), γδ T (CD3−γδ+), and NKT (CD3−CD49b+) cells. (C) Percentage ± SEM of CD4+ or CD8+ cells among the CD45+CD3+ T cell population. (D) Percentage ± SEM of Ly6G−Ly6C+ neutrophils, Ly6G+Ly6C+ monocytic MDSC and F4/80+ macrophages among the CD45+CD11b+ myeloid cell population. Three independent experiments were performed and a representative one is shown. E) ADCC assay was performed using 4T1 target cells incubated with 1:50 pooled sera from vaccinated mice (AX09-0M6, gray bars; MS2 wt, black bars and untreated, white bars) and SPC effector cells at different effector/target cells ratios (200:1, 100:1, and 50:1). Results shown are the mean ± SEM of the percentage of ADCC. **, P < 0.01, ***; P < 0.01, ****; P < 0.001, Student’s t-test.
tumor bulk. Studies investigating combination therapies with AX09-0M6 are being planned.

Targeting self-antigens can raise safety concerns about effecting normal cells and the induction of autoimmune disease. The xCT protein is highly restricted to a few normal cell types, mostly myeloid cells in the spleen and astrocytes and microglia in the central nervous system. We have observed no toxicity when inhibiting xCT function in vivo using AX09-0M6 and DNA-based vaccines. In particular, no inflammatory cell infiltrates or any other abnormalities were detected in brains from both BALB/c and CD-1 mice immunized with AX09-0M6. We believe this result is particularly valuable for the clinical development of AX09-0M6 because it shows that the vaccine generates very robust antibody response in both inbred and outbred mice (e.g. representative of human diversity) with apparently limited or no toxicity.

VLPs targeting both self-antigens and infectious agents (including FDA approved VLP-based vaccines) have demonstrated excellent safety profiles in the clinic. While VLPs elicit high titer antibodies against self-antigens, these antibody responses wane over time. Pharmacokinetic data in human clinical trials performed by Cytos using a similar bacteriophage VLP as AX09-0M6, showed antibody titers against self-antigens had half-lives of ~3 months, which is in stark contrast to long-lived antibody responses using VLP-based vaccines targeting pathogens.\(^{51-53}\) There has been no evidence for antibody boosting by tissue expression of the endogenous self-antigen in animal models, presumably because foreign T-cell helper responses are required to boost antibody production. Therefore, the duration and magnitude of xCT antibodies are controlled by AX09-0M6 dose and dosing schedule, reducing the possibility of autoimmune disease. Also, the xCT octamer (amino acids 443–450) carried by AX09-0M6 is smaller than effective T-cell epitopes, which nearly eliminates the potential of inducing an autoreactive xCT T-cell response. This apparent lack of a CTL response against xCT was endorsed by the inability of SPC from AX09-0M6 vaccinated mice to kill 4T1 cells in vitro and to upregulate CD107a/b expression after 6 days of co-culture with xCT\(^+\) 4T1 cells (data not shown). Nevertheless, the vaccine clearly elicits a T helper 1 response, as indicated by the production of IgG2a against xCT, but this T cell response is against T cell epitopes provided by the VLP proteins that act as carriers.\(^{52,54,55}\) The small size of the xCT epitope carried by AX09-0M6, however, also increases the probability of homology to other proteins and possible off target affects. Indeed, xCT is one member of a large, highly homologous family of heterogeneous amino acid transporters,\(^{56\text{–}58}\) most of which are widely expressed by normal tissues and/or have been reported to be upregulated in a variety of tumors.\(^{57}\) We have blasted the ECD6 peptide with the sequence of most members of the Hetero-meric Aminoacid Transporters family, including LAT1,\(^{58}\) the most highly homologous to xCT, showing that it is not shared. Further protein homology and cellular localization bioinformatics will be used to identify any other potential cross-reactive target.

Ultimately, a VLP-based immunotherapy with no toxicity has a great potential to be used in a maintenance/neo-adjuvant setting for TNBC before tumor recurrence/refractory. AX09-0M6 could slow or stop the cancer’s return by blocking residual CSC that represents a major barrier towards effective cancer treatments.

**Materials and methods**

**Cell and tumorsphere generation**

4T1 and HCC-1806 cells were grown in RPMI-1640 Medium (Invitrogen, Corp.) supplemented with 10% heat-inactivated Fetal Bovine Serum (FBS; Sigma-Aldrich). MDA-MB-231 cells

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**Figure 7.** AX09-0M6 is efficacious in preventative and therapeutic models of aggressive breast cancer. (A) Passage 1 tumorspheres derived from 4T1 cells were injected into the inguinal mammary fat pad. When the tumors reached 1 – 1.5 mm in diameter, mice were treated with MS2 wt and AX09-0M6, and boosted 14-days later. Primary tumor diameters were measured at the indicated time-points and tumor volume was calculated. Results shown are the average tumor volume from two independent experiments and the error bars represent the standard deviation of the mean. AX09-0M6 vs MS2 wt: *P < 0.05; **P < 0.005; Student’s t-test. AX09-0M6 vs untreated: P < 0.005 from week 26 to 33; P < 0.05 from week 35; Student’s t-test. (B) Mice from A were euthanized 40 days after tumor challenge and the number of micrometastases was enumerated. Each dot represents a single animal and is the average number of lung metastases from at least 2 sections. *P = 0.0126; **P = 0.0046, Mann Whitney test. Results shown are the aggregate from 2 independent experiments.
were grown in Dulbecco’s Modified Eagle Medium (DMEM, Invitrogen Corp.) supplemented with 10% FBS. All cells were purchased from ATCC (LGC Standards). TUBO cells were cultured in DMEM supplemented with 20% FBS. Each culture medium was supplemented with Penicillin (100 U/mL) and Streptomycin (100 U/mL) (Sigma-Aldrich). Cells were cultured at 37°C in a 5% CO₂ incubator and all cells were tested negative for mycoplasma using polymerase chain reaction (PCR) MycoAlert assay (Lonza). Passage 1 and passage 2 (P1 and P2) tumorspheres were generated as previously described.

Production and purification of VLPs
Production and purification of single-chain dimer versions of MS2 bacteriophage VLPs were done as described.Briefly, ECD6 peptide sequence (LYSDPFST) derived from the sixth loop of XCT protein was inserted genetically into the AB loop of MS2 CP through pDSP62 plasmid as previously described. Plasmids were electroporated into the E. coli, T7 expression strain C41 (DE3) (Lucigen), grown to mid-log phase and VLP expression was induced by the addition of IPTG (1 mM, Sigma-Aldrich) overnight at 37°C. Bacteria pellets were lysed in lysis buffer (50 mM Tris-HCl, pH 8.5, 100 mM NaCl, 10 mM EDTA), sonicated and purified from bacterial debris by centrifugation. Bacterial DNA was removed by treating the supernatant with DNaseI (10 units/mL, 1 hour at 37°C; Sigma-Aldrich) and the VLPs were purified by size exclusion chromatography (Sepharose CL-4B resin, Sigma-Aldrich). Buffer exchange (Phosphate-buffered saline, PBS) and concentration of purified VLPs was achieved using ultrafiltration (Amicon Ultra device, 100Kd MWCO) and resulting VLP preparations were quantitated by Bradford assay (BioRad). VLP purity was assessed by agarose and SDS-PAGE gel electrophoresis.

Purification of IgG for in vitro assays
Female BALB/c mice were administrated 10 µg of VLP without exogenous adjuvant by intramuscular (i.m.) injection followed by a single boost 28 days later. Two weeks after final boost, blood was collected by cardiocentesis and immune sera were isolated as previously described. Total IgG were purified by size exclusion chromatography (Phasarope CL-4B resin, Sigma-Aldrich). Buffer exchange (Phosphate-buffered saline, PBS) and concentration of purified IgG was achieved using ultrafiltration (Amicon Ultrafiltration device, 100Kd MWCO) and resulting VLP preparations were quantitated by Bradford assay (BioRad). VLP purity was assessed by agarose and SDS-PAGE gel electrophoresis.

Characterization of antibody responses
XCT specific antibody responses were evaluated by ELISA using ECD6 peptide (synthesized with a CGG linker at the C-terminus; GenScript) as coating antigen. Avidin (Pierce) prepared in PBS pH 7.5 was added to wells of Immulon 2HB ELISA Plates (Thermo Scientific) at 500 ng/well and incubated for 2 h at 37°C. Following washing, SMPH (Succinimidyl 6-((beta-maleimidopropionamido)hexanoate); Thermo Scientific) was added to wells at 1 µg/well and incubated for 2 h at 37°C. ECD6 peptide was added at 1 µg/well and incubated overnight at 4°C. Plates were blocked with 0.5% milk in PBS at room temperature for 1 h, and 3-fold dilutions (starting at 1:500) of sera from individual animals were added to each well and incubated for 2 h. Wells were probed with HRP-labeled goat anti-mouse IgG (Invitrogen; 1:5000) for 1 h. The reaction was developed with TMB Solution (3’,3’,5’,5’-Tetramethylbenzidine, Calbiochem) for 1 h and the reaction was stopped using 1% HCl. ECD6 activity was determined by measuring optical density (OD) at 450 nm. Wells with twice the OD450 value of background (sera from MS2 wt mice) were considered positive and the highest dilution with a positive value was considered the endpoint dilution titer.

To determine IgG subclass elicited by VLP administration, ELISA plates were coated as above. Total IgG and each IgG subclass were measured in triplicate from each animal using a sera dilution of 1:500. Wells were probed with HRP-labeled goat anti-mouse IgG, IgG1, IgG2a, and IgG2b (Invitrogen; 1:5000) for 1 h. Detection was as described above. Relative ratio of each IgG subclass was calculated by taking the average OD450 value from IgGx divided by the average OD450 from total IgG.

Immune sera effect on tumorspheres
P1 tumorspheres were dissociated and cultured at a density of 6 × 10⁴ cells/ml in a 6-well dish in the absence or presence of 50 µg/ml of purified IgG or of 50 µM SASP (sulfasalazine; Sigma-Aldrich). After 5 days P2 tumorspheres were imaged with an ApoTome fluorescence microscope (Zeiss) and sphere diameter measured with the AxioVision 4.8 software as described. Spheres were counted and reported as number of spheres generated for every 10³ cells plated, then they were dissociated and processed for FACS analysis of Aldefluor and ROS.

Immunofluorescence
P2 tumorspheres were cytopspun to glass slides, fixed in 4% formalin solution and then blocked in 10% bovine serum albumin (BSA, Sigma-Aldrich). Slides were incubated with 100 µg/ml of purified IgG or 1:10 dilution of sera from CD-1 mice for one hour at room temperature, washed with PBS and incubated with rabbit AlexaFluor488 anti-mouse (Life Technologies). Nuclei were stained with DAPI (Sigma-Aldrich). Images were acquired with a Leica TCS-SP5 II confocal microscope and analyzed using LASAF software (Leica). Fluorescence was quantitated using ImageJ and signal was calculated as integrated density normalized to nuclei number.

In vivo treatments
Female 6- to 8-week-old BALB/c mice (Charles River Laboratories Italia) were maintained at the Molecular Biotechnology Center, University of Torino, and treated in accordance with the University Ethical Committee and European guidelines under Directive 2010/63. Vaccination, performed either before (preventive model) or after tumor challenge (therapeutic model), consisted of two i.m. injections at 2 week intervals. Each mouse
received 5-10 μg of MS2 wt or AX09-0M6, formulated in PBS for a total volume of 50 μL. In the preventive model, one week after last vaccination 5 × 10⁴ tumorspheres derived from TUBO cells were injected i.v. Mice were sacrificed at 20 days after cell challenge. In the therapeutic model, 1 × 10⁴ P1 4T1 tumorsphere derived cells were injected s.c. and when the tumor reached a mean diameter of 1-1.5 mm, mice were vaccinated as described above. Tumors were then measured twice a week with calipers in two perpendicular diameters. Tumor growth was reported as volume calculated as \((\pi/6) \times A^2 \times B\) where A and B represent the short and long diameters, respectively. Mice were euthanized 40 days after tumor cell challenge. At sacrifice, blood was collected by cardiocentesis and sera were stored at -20°C for use in immunoassays. Lungs were removed, the right lung was fixed in formaldehyde solution (4%), paraffin embedded, sectioned, and Hematoxylin and Eosin (H&E) stained. Micrometastases were counted with a Nikon SMZ1000 stereomicroscope and analyzed using ImageJ software (U.S. National Institutes of Health).

**FACS analysis**

Tumorsphere derived cells were stained with Aldefluor kit (Stem Cell Technologies) as reported. To measure intracellular ROS content, cells were stained with 2’,7’-dihydrochlorofluorescein diacetate (DHCF-DA, Sigma-Aldrich) as described. To quantify anti-xCT antibody titers, 5 × 10⁴ tumorsphere-derived cells were incubated for 30 min at 4°C with 50 μg/mL of IgG purified from sera of vaccinated mice. Cells were pre-treated with Fc receptor (FcR) blocker (CD16/CD32; Becton Dickinson) for 5 min at 4°C. After washes with PBS containing 2% BSA (Sigma-Aldrich) and 0.1% sodium azide (NaN₃; Sigma-Aldrich), cells were incubated with a 1:50 dilution of a FITC-conjugated anti-mouse IgG antibody (DakoCytomation) for 30 min at 4°C. Rabbit anti-xCT Ab (Thermo Fisher Scientific) followed by FITC-conjugated anti-rabbit IgG Ab (DakoCytomation) was used as a positive control on cells fixed and permeabilized with BD Cytofix/Cytoperm kit (Becton Dickinson). Washed cells were then acquired on the BD FACSVerse and analyzed with BD FACSuite™ software (Becton Dickinson). The results were expressed as percentage of positive cells and as mean fluorescence intensity (MFI).

For the analysis of immune infiltrates, lungs from vaccinated mice were finely minced with scissors and then digested by incubation with 1 mg/ml collagenase IV (Sigma Aldrich) in RPMI-1640 (Life Technologies) at 37°C for 1 h in an orbital shaker. After washing in PBS supplemented with 2% FBS, the cell suspension was incubated with a buffer for erythrocyte lysis (155 mM NH₄Cl, 15.8 mM Na₂CO₃, 1 mM EDTA, pH 7.3) for 10 minutes at R.T., then washed through a 70-μm pore cell strainer, centrifuged at 1400 rpm for 10 min and re-suspended in PBS, treated with Fc receptor blocker as described. Cells were then stained with the following Abs: anti-mouse CD45 VioGreen, anti-mouse CD3 FITC, anti-mouse CD4 APC-Vio770, anti-mouse CD8 VioBlue, anti-mouse γδ PE/Vio770, anti-mouse CD49b PE, anti-mouse F4/80 PE/Vio770, anti-mouse CD11b FITC, anti-mouse Ly6G Vioblue, and anti-mouse Ly6C APC/Vio770 (all from Miltenyi Biotec). For the analysis of xCT expression on myeloid SPC, spleens from vaccinated mice were smashed on a 70-μm pore cell strainer, erythrocytes were lysed, FcR blocker and cells were stained with anti-mouse CD45 VioGreen, anti-mouse F4/80 PE/Vio770, anti-mouse Ly6G Vioblue, anti-mouse Ly6C APC/Vio770, anti-mouse class II HLA APC (Miltenyi Biotec) and anti-mouse CD206 PE (ebiosis). After washing, cells were fixed and permeabilized with BD Cytofix/Cytoperm kit and stained with rabbit anti-xCT Ab followed by FITC-conjugated anti-rabbit IgG Ab. Samples were acquired on a BD FACSVerse and analyzed using the BD FACSuite software.

**Antibody-dependent cell-mediated cytotoxicity (ADCC)**

1 × 10⁴ 4T1 target cells were labeled with 2 μM of carboxyfluorescein diacetate succinimidyl ester (CFSE; Molecular Probes) following the manufacturer’s instructions. ADCC was initiated by the addition of SPC from untreated BALB/c mice as effector cells (200:1, 100:1, and 50:1 as the effector: target (E:T) ratio) and sera from vaccinated mice at a 1:50 dilution. The plate was incubated overnight at 37°C, then cells were harvested, stained with 1 μg/ml 7-Amino-Actinomycin D (7-AAD, BD Bioscience), acquired on the BD FACSVerse and analyzed by BD FACSuite™ software (Becton Dickinson). Percent killing was obtained by back-gating on the CFSE⁺ targets and measuring the % of 7-AAD⁺ dead cells. % ADCC for each serum sample was calculated with the formula \[\text{ADCC} = \left(\frac{\text{dead targets in sample} – \text{spontaneously dead targets}}{\text{dead target maximum - spontaneously dead targets}}\right) \times 100.\] Spontaneous release was obtained by incubating target cells in medium with serum, whereas maximal release was obtained after treatment with triton solution.

**Histological and immunohistochemical analysis**

Mice were killed by anaesthesia overdose, and explanted brains were fixed in 4% formalin for 24 hours and then sliced in coronal sections and processed for histology. Paraffin sections were cut at 5 μm and stained with Hematoxylin and Eosin (H&E) or with immunohistochemical antibodies. For the immunohistochemical detection of microglial cells, non-specific immunoreactivity was blocked by incubation of the brain sections with 5% goat serum and 3% milk. The slides were incubated for 1 hour with a rat anti-mouse F4/80 primary antibody (Abcam, CI:A3-1). Biotinylated anti-rat secondary antibody (Dako) was used and the avidin-biotin-peroxidase complex (ABC reaction, Vector Lab) was visualized with diaminobenzidine (Thermo Scientific). Sections were counterstained with haematoxylin before dehydration and mounting. Slides were digitized with the Panoramic Desk scanner and subsequently evaluated by a pathologist with Panoramic Viewer (both from 3D HISTECH).

**Statistical analysis**

Differences in sphere formation, protein expression, number of infiltrating cells and proliferation index were evaluated by a Student’s t-test. Differences in number of metastases were evaluated by a non-parametric Mann Whitney test. Data are shown as mean ± SEM unless otherwise stated. Values of \(P < 0.05\) were considered significant.
Conflict of interest
The authors declare no potential conflicts of interest.

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