An alphavirus vector overcomes the presence of neutralizing antibodies and elevated numbers of Tregs to induce immune responses in humans with advanced cancer

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Therapeutic anticancer vaccines are designed to boost patients’ immune responses to tumors. One approach is to use a viral vector to deliver antigen to in situ DCs, which then activate tumor-specific T cell and antibody responses. However, vector-specific neutralizing antibodies and suppressive cell populations such as Tregs remain great challenges to the efficacy of this approach. We report here that an alphavirus vector, packaged in virus-like replicon particles (VRP) and capable of efficiently infecting DCs, could be repeatedly administered to patients with metastatic cancer expressing the tumor antigen carcinoembryonic antigen (CEA) and that it overcame high titers of neutralizing antibodies and elevated Treg levels to induce clinically relevant CEA-specific T cell and antibody responses. The CEA-specific antibodies mediated antibody-dependent cellular cytotoxicity against tumor cells from human colorectal cancer metastases. In addition, patients with CEA-specific T cell responses exhibited longer overall survival. These data suggest that VRP-based vectors can overcome the presence of neutralizing antibodies to break tolerance to self-antigen and may be clinically useful for immunotherapy in the setting of tumor-induced immunosuppression.

Introduction

Efficacious therapeutic anticancer vaccines are now becoming a reality, but a number of challenges remain. More efficient platforms for delivering antigen to in situ DCs, critical for activation of T cell and antibody responses (1), are needed. Viral vectors are particularly promising for tumor antigen delivery because many can directly infect DCs and, by the nature of their engagement of innate immune signals, provide the appropriate cognate costimulation or “danger signals” that lead to enhanced DC activation breaking of tolerance through induction as well as expansion of the adaptive immune response. There has been extensive experience with numerous viral vectors based on poxvirus (vaccinia, fowlpox, canarypox), adenovirus, reovirus, herpesvirus, and others. Nonetheless, a challenge to the use of all viral vectors is the induction of neutralizing antibodies to proteins on the virus surface that precludes the repeated administration of recombinant vector-based vaccines, particularly those based on vaccinia and adenovirus (2, 3).

An attractive alternative is vaccination with recombinant alphavirus such as Venezuelan equine encephalitis (VEE) virus. The VEE structural protein genes may be replaced with a heterologous gene of interest to produce a recombinant RNA replicon capable of self replication and very high-level expression of the foreign gene, often reaching levels of 10%–15% of the total cell protein. These replicons may be packaged into virus-like replicon particles (VRP) by cotransfection of cells in vitro with helper RNAs encoding the viral structural proteins (4). VRP are morphologically identical to alphaviruses and efficiently express their foreign gene(s) at the site of inoculation, but the absence of the structural protein genes on the replicon prevents further amplification in the vaccine recipient. Other advantages of VRP are their tropism for, and maturation of (5), DCs, which could result in enhanced T cell activation. A number of preclinical studies have reported that VRP induce potent immunity despite the presence of neutralizing antibodies (4–12).

Recently, an alphavirus replicon vaccine for CMV was tested in CMV-seronegative volunteers. The vaccine was well tolerated and induced T cell and antibody responses against the highly immunogenic CMV proteins in these healthy young subjects (13). A critical question was whether we could immunize multiple times with the same vector against a self antigen if anti-VRP neutralizing antibodies were induced. We therefore tested to determine whether a VRP-based vector could be repeatedly administered to break tolerance to a nonimmunogenic self antigen in a clinically relevant setting, such as the immunomodulated environment of older patients with metastatic cancer.

We chose as the tumor antigen carcinoembryonic antigen (CEA) (14), widely expressed in colon, breast, lung, and other malignancies and well established as an immunotherapy target for cancer vaccines (15). We designed a VRP vaccine expressing CEA(6D), an Asn to Asp substitution in the CEA gene within the peptide capable of binding to HLA-A2, resulting in enhanced recognition by the T cell receptor (16). We tested its immunogenicity in murine models and demonstrated our ability to achieve improved immune response in preclinical models with a prime-and-boost strategy. We then gener-
ated GMP grade vector for this phase I/II clinical trial to evaluate the safety and clinical and immunologic efficacy of immunization with CEA(6D)-VRP (AVX701) in patients with advanced cancer. The use of any cancer vaccine strategy must address the reality that there are a number of immunoregulatory mechanisms, in particular, Tregs, invoked in the tumor microenvironment, regional lymph nodes, and peripheral blood that may limit or prevent the induction of clinically effective adaptive immune responses (17). CD4 \(^{pos} \) CD25 \(^{hi} \) FoxP3 \(^{pos} \) Tregs prevent uncontrolled proliferation of antigen-specific T cells (18). Elevated Treg levels can be detected in the peripheral blood, regional lymph nodes, and the tumor microenvironment of cancer patients compared with healthy individuals and are negatively associated with survival (19). Because Tregs appear to impair host immune responses against cancer (20), one approach to eliminating the influence of Tregs is by depleting them with the CD25-targeting immunotoxin denileukin diftitox prior to immunization (as others and we have done) (21, 22) or anti-CD25 antibody. However, these approaches could interfere with activated T cells that also express CD25. Viral vectors have been reported to provide persistent Toll-like receptor signals capable of reversing Treg-mediated CD8 tolerance in murine models (23). Therefore, we were interested in determining whether VRP-CEA(6D), would induce an adaptive immune response to self antigen in the setting of an immunosuppressive environment caused by elevated Tregs.

We performed a traditional “3 + 3” phase I study of active immunotherapy with VRP-CEA(6D). We analyzed blood samples before each immunization and following completion of all immunizations for VRP-neutralizing antibodies, circulating Treg and CEA-specific adaptive humoral and cellular immune responses. Although we noted elevated levels of Tregs in the study patients and observed a rapid induction of anti-VRP neutralizing antibodies following the first immunization, we found that potent CEA-specific T cell and antibody responses could nonetheless be induced by VRP-CEA(6D).

Results

Choice of multiple immunizations. We wished to develop an alphavirus-based vaccine in order to take advantage of the alphavirus’ tropism for DCs and ability to generate large amounts of a protein of interest after infecting a cell. This was expected to cause infected DCs at the injection site to process and appropriately present CEA tumor antigen. Prior to initiating the clinical trial, we determined whether a single dose or a prime-plus-boost dose of VRP-CEA would achieve the highest levels of T cell and antibody responses. Although we noted elevated levels of Tregs in the study patients and observed a rapid induction of anti-VRP neutralizing antibodies following the first immunization, we found that potent CEA-specific T cell and antibody responses could nonetheless be induced by VRP-CEA(6D).

Table 1

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Notes: Numbers in parentheses show prior CEA vaccine treatments. All patients had stage 4 cancer; those with 0 sites of metastases had undergone resection of their tumor or had no evidence of disease after chemotherapy. KPS, Karnofsky performance status; Ph., phase; NSCLC, non–small cell lung cancer.
addition, therapeutic antitumor effects had been demonstrated in models following multiple vaccinations (data not shown). Furthermore, clinical trials targeting CEA have used multiple injections or priming followed by heterologous boosting. We therefore chose multiple immunizations for the clinical trial testing of this vector in patients with advanced cancer.

**Patient demographics.** Patients (18 male, 10 female, ages 36–69) who were diagnosed with colorectal (n = 23), appendiceal (n = 1), pancreas (n = 1), lung (n = 2), or breast (n = 1) cancer, had a performance status of 80%–100%, and had failed a median of 4 prior chemotherapeutic regimens (range 2–11) were entered into the trial by cohorts and treated at 1 of 3 dose levels (Table 1). In the first cohort, 3 of 4 patients completed all 4 immunizations. One dropped out of the study due to development of a brain metastasis. In the second cohort, 3 of 3 patients completed all immunizations. In the third cohort, 6 of 7 patients completed all immunizations (with 1 dropping out after 3 immunizations due to progressive disease) and 2 continued receiving additional booster doses afterwards. Dose-limiting toxicity (DLT) was not reached, and the dose of 4 × 10^8 IU was determined to be the maximal feasible dose. The trial allowed for 14 additional patients to be treated at the maximal feasible dose (4 × 10^8 IU). Of these next 14 patients, 10 received at least 4 immunizations, and 4 discontinued prior to the final immunization, all due to progression of disease. Two received an additional booster immunization. Overall, immunizations were well tolerated with 6 grade 3 events (1 each of alanine aminotransferase [ALT], bilirubin, neurological confusion, obstruction, dyspnea, and fatigue, all attributed to disease progression) and no grade 4 or higher events. The only injection site reactions observed were grade 2 instances of injection site pain. There were no febrile reactions.

**Elevated levels of Tregs in patients.** We demonstrated that our gating strategy detecting CD4^+CD25^+ cells contained high levels of FoxP3 and could be sorted into a population that had immunosuppressive effects in vitro (Supplemental Figure 2). We compared the circulating CD4^posCD25^posFoxP3^pos Treg levels in the patients participating in this study with those of healthy volunteers. Consistent with previous reports (17), the percentage of Tregs was statistically greater (P = 0.03) for patients than healthy controls (Figure 1A). There was no statistically significant increase in Tregs following immunization both by percentage of CD4^+ T cells and absolute number of CD4^posCD25^posFoxP3^pos Tregs per microliter of blood (Figure 1, A and B).

Antigen-specific T cell and antibody responses despite neutralizing antibody and elevated Treg levels. We measured the induction of anti-VRP neutralizing antibodies in patients in the high-dose cohort. We observed, after a single dose of VRP-CEA(6D), that the anti-VRP antibody titer increased dramatically and remained elevated throughout the immunizations (Figure 2A). Titors also remained elevated in the 4 patients in the high-dose cohort who continued to receive booster vaccinations every 3 months (with follow-up reaching 12 months) but not in the lower-dose cohorts (Supplemental Figure 3). Because we measured the immune response prior to each injection, we observed that the T cell response progressively increased beginning with the second or third immunization (Figure 2, B and C). For the 4 patients who received ongoing booster doses, the immune response levels remained at the levels induced by the initial 4 doses (data not shown). Again, we did not have long-term specimens for patients who received only the first 4 immunizations; however, because the first booster occurred at 3 months following the fourth injection, the data do suggest that the anti-VRP titers remain elevated at least 3 months following the first series of 4 injections.

The CEA-specific immune responses were analyzed at each time point by ELISpot and ELISA (Supplemental Figure 3 and Figure 2, B and C). We found a significant increase in immune response for at least 1 postvaccination time point versus prevaccination for all 3 assays in the high-dose cohort but not in the lower-dose cohorts (Supplemental Figure 3). Because we measured the immune response prior to each injection, we observed that the T cell response progressively increased beginning with the second or third immunization (Figure 2, B and C). For the 4 patients who received ongoing booster doses, the immune response levels remained at the levels induced by the initial 4 doses (data not shown). Again, we did not have long-term specimens for patients who received only the first 4 immunizations; however, because the first booster was at 3 months following the fourth injection, the data do suggest that the CEA-specific immune responses remain elevated at least 3 months following the first series of 4 injections. The data also suggest that further boosting does not continue to enhance the immune response, perhaps because the maximum attainable response has been reached.

We further characterized the ELISpot results using intracellular cytokine staining to identify the CD4^+ (Figure 2D), and CD8^+ CEA-specific T cell responses. The CD4^+ CEA-specific T cell response increased by the final immunization in the majority (9/13) of the patients. A CEA-specific CD8^+ T cell response was also observed in 5/9 patients analyzed in the high-dose cohort (data not shown). Taken together, these results illustrate that VRP-CEA(6D) induces both CD4^+ and CD8^+ CEA-specific T cell and antibody responses.

We then determined whether patients with low Tregs, defined as below the median of the Treg level for all patients in the study, had a different magnitude of immune response than those with high Tregs (i.e., above the median of the Treg levels for all patients).
Figure 2
Immune analysis following VRP-CEA(6D) vaccination. Patient sera or PBMCs were analyzed before and after VRP-CEA(6D) for each immunization for MTD cohort by (A) anti-VRP microneutralization assay; (B) CEA ELISA; (C) VRP-CEA ELISpot; and (D) intracellular IFN-γ. (A) Patient sera were analyzed for anti-VRP antibodies by microneutralization assay for weeks 0, 3, 6, 9, and 12. The antibody titer is presented for each week as mean ± SD. The endpoint titer was defined as the last serum dilution at which there was at least 80% reduction in the number of GFP-positive cells compared with control wells. (B) Patient sera were tested against CEA protein by ELISA for weeks 0, 3, 6, 9, and 12. The antibody titer is represented over the course of the 4 vaccinations as mean ± SEM. A single patient with an outlier titer of 1:1,600 at week 9 was not included in the data presented in the graph but was included for statistical analysis. *P = 0.02. (C) Patient PBMCs were stimulated with VRP-CEA (MOI 10) in an ELISpot assay. The number of IFN-γ-producing cells per 10⁶ PBMCs is presented as mean ± SEM. *P = 0.01; **P = 0.005. (D) Patient PBMCs were stimulated with VRP-CEA (MOI 1) and incubated for 12 hours in an intracellular cytokine assay. The percentage of cells that were CD4⁺CD69⁺IFN-γ⁺ is presented as mean ± SEM. Time of vaccinations is represented by arrows on the x axis of graph (D). Regression analysis with repeated measures was used to analyze assay results over time. Time points where the mean significantly differed from baseline are indicated on the graph (P values are adjusted for multiple comparisons). *P = 0.01.

The magnitude of the immune (CEA-specific T cell and antibody) responses was not statistically different regardless of whether patients had low- or high- peripheral blood Treg levels (Figure 3, A–C). Similarly, there was no significant correlation between the magnitude of anti-VRP neutralizing antibody titers and anti-CEA antibody titers or ELISpot levels (Figure 3, D and E). Hence, despite the induction of strong neutralizing antibodies and the presence of elevated Treg levels, the VRP-CEA(6D) remained capable of activating potent anti-CEA antibody and T cell responses.

Antitumor activity of vaccine-induced anti-CEA antibodies. We wished to determine the functional activity of vaccine-induced anti-CEA antibodies by analyzing whether antibodies in patient serum could bind a CEA-expressing colon cancer cell line and mediate antibody-dependent cellular cytotoxicity (ADCC) (Figure 4). CEA-specific antibodies in the sera of 2 patients from the high-dose cohort bound to the CEA⁺ cell line (SW1463) but not to the CEA⁺ cell line (COLO320) at titers of 1:50 and 1:100 (Figure 4A). Furthermore, the CEA-specific antibody sera could mediate ADCC (Figure 4B). These data confirm that antibodies induced by VRP-CEA(6D) have functional capacity despite the presence of anti-VRP neutralizing antibodies and high Treg levels.

Antibodies developed after a single injection and peripheral blood Tregs were elevated in a substantial fraction of these advanced cancer patients, yet the VRP-CEA induced similarly high level CEA-specific T cell and antibody responses with repeated doses of the same vector (homologous prime boost).

Although immunization against tumor antigens has been achieved with heterologous prime-boost strategies, homologous prime-boost strategies with vaccinia or adenovirus vectors have previously had limited success due to the induction of neutralizing antibodies that nullify the effect of successive immunizations (2, 3). The precise mechanism for retained functionality of VRP-based vaccines despite antivector neutralization is not known. However, a possible explanation could be the origin of the strain of VEE used for generating the glycoprotein coat of the VRP. The strain, named V3014, was generated following passages under a selective pressure for accelerated penetration (24, 25) that resulted in multiple genetic alterations. It is possible that this rapidly cell-penetrating phenotype rendered the virion less sensitive to neutralization by virtue of being able to bind and infect cells in vivo before effective neutralization occurs. Another explanation, which does not exclude the first, could be the ability of

Clinical outcome. There were 2 patients with stable disease and 1 with a complete response (CR) of a small liver lesion (Figure 5A). Two patients had no evidence of disease prior to immunization and remained in that condition. Although numbers were small and there were not enough events to make a statistical comparison, nonetheless, there appeared to be better survival in the patients with an immune response by ELISpot (Figure 5B).

Discussion
The purpose of this study was to determine whether a vaccine based on recombinant alphaviruses could be repeatedly administered to break tolerance to a self antigen despite the expected elevated levels of immunosuppressive Tregs and the induction of neutralizing antibody against the VRP. Notwithstanding these challenges to vaccination, we hypothesized we would be able to break tolerance because preclinical studies had suggested higher magnitudes of T cell response after multiple immunizations with VRP-CEA compared with a single immunization (Supplemental Figure 1). Furthermore, in other preclinical studies (Supplemental Figure 4), we observed that even when VRP-encoding antigen was mixed with enough anti-VRP antibody to achieve neutralization of the VRP and then injected, there was still a high rate of T cell and antibody response against the antigen. Indeed, in the present clinical study, we observed that neutralizing
VRP to target cells efficiently in vivo even when coated with otherwise neutralizing antibodies. Another explanation could be the fact that VRP is able to infect and replicate its genome in DCs, which are capable of taking up exogenous particular material such as opsonized virus. Finally, because VRP generate large amounts of the encoded protein (CEA), even a small number of VRP reaching DCs without neutralizing antibody could be sufficient to induce potent immune responses.

There are several possible explanations for why we observed similar magnitudes and rates of CEA-specific T cell and antibody responses regardless of the level of Tregs. Viral vectors have been reported to provide persistent Toll-like receptor signals capable of reversing Treg-mediated CD8 tolerance in murine models (23). Of course, it is possible that the Foxp3⁺ cells are not functional Tregs or that the actual anatomic location for breaking Treg-mediated tolerance to antibody and T cell response is not blood. However, we have previously observed that blood Tregs are functional (Supplemental Figure 2) and depleting them from peripheral blood was associated with enhanced T cell responses (21). Therefore, we believe that the elevated Tregs observed in the peripheral blood of the patients in the current study would have the capability of inhibiting T cell responses, yet the alphavirus vaccine was able to evade this effect. There is less published on the effect of Tregs on antibody responses, however.

Fields reported that the secretion of antibodies by anti-chromatin B cells is blocked by the injection of CD4⁺CD25⁺FoxP3⁺ Treg cells (26). This suggests that Tregs can have a negative effect on antibody responses as well.

Although the rate of clinical responses was low in this heavily pretreated population, we did observe a regression of a liver metastasis in a patient with pancreatic cancer; 2 patients had stable disease, and 2 patients with no evidence of disease remained in remission. All of these instances of clinical benefit were in the high-dose cohort, suggesting the possibility of a dose–clinical response relationship. It is possible we would see greater clinical benefit with higher doses of VRP-CEA. However, for what we believe is the first-in-human clinical study, we limited the dose to 4 × 10⁸ IU, which was the highest level for which we had preclinical toxicology data available.

In summary, we were able to treat advanced cancer patients with repeated immunizations with a recombinant alphaviral vector and to break tolerance to nonmutated CEA despite the induction of neutralizing antibody against the VRP and elevated Treg levels. This is the first study we are aware of that shows a viral vector overcoming neutralizing antibodies and elevated Tregs in cancer patients. Future studies will focus on enhancing the immune response further by the addition of cytokines such as IL-12 and studying the VRP in patients with less advanced disease.
Methods

**Generation of CEA(6D) VRP.** We generated a replicon expression plasmid containing the CEA gene with the Cap-1(6D) mutation (15) called CEA(6D) VRP (AVX701) (Supplemental Figure 5) as previously described (9, 13, 27). It was formulated so that the maximum feasible dose to administer in the clinical trial was \(4 \times 10^8\) IU.

**Preclinical studies to assess immunogenicity in mice.** Human-CEA transgenic mice (28) were vaccinated via bilateral footpad injection on day 0 with VRP-CEA \((1 \times 10^7\) virus particles) and day 21 with either saline or VRP-CEA. Fourteen days later, mice were euthanized and splenocytes were collected for analysis. IFN-\(\gamma\) ELISpot assay was performed to examine the CEA-specific T cell responses (4 mice per condition) to an overlapping polypeptide mix for human CEA. All animal studies were approved by the Duke IACUC.

**Protocol schema and patient treatment.** The study was performed under an FDA-approved Investigational New Drug Exemption (IND) and registered at ClinicalTrials.gov (NCT00529984). Participants were recruited from oncology clinics at Duke University Medical Center and provided signed informed consent approved by the Duke Institutional Review Board. Eligibility requirements included metastatic cancer expressing CEA and adequate hematologic, renal, and hepatic function. Trial participants were required to have received treatment with standard therapy known to have a possible overall survival benefit or refused such therapy. Exclusion criteria included chemotherapy or radiation within the prior 4 weeks, history of autoimmune disease, viral hepatitis, HIV, or use of immunosuppressives. Some patients continued bevacizumab or cetuximab. Prior CEA immunotherapy was permitted.

This study used a standard 3 + 3 dose escalation strategy with DLTs defined as grade 3 or 4 major organ toxicity. The CEA(6D) VRP injections were given....

**Figure 4**

Serum cell binding and ADCC assay. (A) Sera from high-dose cohort patient AVX701-0121 were incubated with CEA-expressing colon tumor cell line SW1463 (CEA+) and non-CEA-expressing colon tumor line COLO320 (CEA-). Results are represented as histograms of MFI for 1:50, 1:100, and 1:200 dilutions. Binding of prevaccination serum antibodies is represented by white peaks, and the postvaccination serum is represented by the gray peaks. The top row of histograms binds against SW1463 (CEA+) cell line and the lower row COLO320 cell line. The ELISA titer for patient AVX701-0121 was 1:200. (B) Serial dilutions of pre- and postvaccination sera from high-dose cohort patient AVX701-0109 were incubated with CEA-expressing colon tumor cell line SW1463 (CEA+) in an ADCC assay. The percentage of specific lysis is presented as mean ± SD. *\(P < 0.01\) for prevaccination versus week 9 serum at 1:20 ratio. The ELISA titer for patient AVX701-0109 was 1:1,600.

**Figure 5**

Clinical response. (A) CT scan from a pancreatic cancer patient refractory to chemotherapy and radiotherapy. Arrow indicates a liver metastasis that resolved after vaccination. (B) Kaplan-Meier estimates of overall survival by best immune response (by ELISpot) for patients treated on VRP-CEA(6D) (AVX701) at MTD \((4 \times 10^8\) IU).
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into the deltid muscle and alternated side-to-side at each visit. The doses were as follows: cohort 1: CEA(6D) VRP at a dose of 4 × 10⁷ IU in 0.5 ml i.m. every 3 weeks for 4 immunizations; cohort 2: CEA(6D) VRP at a dose of 1 × 10⁸ IU in 0.125 ml i.m. every 3 weeks for 4 immunizations; cohort 3: CEA(6D) VRP at a dose of 4 × 10⁸ IU in 0.5 ml i.m. every 3 weeks for 4 immunizations. Higher doses were not possible due to limits on the amount of clinical-grade study drug. Following establishment of the safety of the dose of 4 × 10⁷ IU, an additional 14 patients received CEA(6D) VRP at this dose. Patients with stable disease after the 4 study injections were permitted to receive booster doses of CEA(6D) VRP every 3 months until progressive disease.

Enumeration of Tregs. Whole blood from before and after immunizations was stained using anti-CD25, anti-CD4, anti-CD3, and anti-CD45 with BD Bioscience TruCount tubes as previously described (ref. 21 and Supplemental Figure 3). CD4⁺CD25⁺ FoxP3⁺ T cells were identified by gating on the CD4⁺CD25⁺ cells, and the percentage of FoxP3⁺ cells and the number of CD4⁺CD25⁺ FoxP3⁺ cells per microliter of blood determined. Peripheral blood was obtained from healthy donors who signed informed consent forms approved by the Duke Institutional Review Board.

Analysis of CEA-specific T cell responses by IFN-γ ELISpot assay and intracellular cytokine staining. For immunofluorescence, freshly isolated PBMCs from peripheral blood were collected at weeks 0, 3, 6, 9, and 12 and analyzed for antigen-specific reactivity by ELISpot and intracellular cytokine assays as previously described (21).

Analysis of anti-CEA antibodies by ELISA. Patient sera were collected at weeks 0, 3, 6, 9, and 12. 96-well plates were coated with whole CEA protein (100 ng/well) and incubated with 100 μl of serum serially diluted 1:10 to 1:31,250. Titers were defined as the highest dilution such that the mean absorbance was equal to twice the negative control.

Analysis of anti-vector responses with a VRP microneutralization assay. To determine anti-vector responses, antibodies to VRP were measured using a microneutralization assay as previously described (13). VRP expressing GFP was mixed with serial dilutions of patient sera and then added to vero cells. The number of cells expressing GFP in each well was determined by fluorescence microscopy.

Analysis of anti-CEA antibody binding to CEA human colon cancer cells by FACS. We have adapted a methodology reported by Wei et al. to measure vaccine-induced antibodies in serum by flow cytometry (29). Serial dilutions of sera were incubated with CEA-expressing colon tumor cell line SW1463 (CEA⁺) and non-CEA-expressing colon tumor line COLO320 (CEA⁻) (3 × 10⁵ cells) for 1 hour at 4°C and analyzed for antibody binding to cells by FACS on a BD LSRII.

ADCC assay. Patient PBMCs were incubated in rHL-2 (1000 IU/ml) overnight. SW1463 (CEA⁺) and Colo320 (CEA⁻) colorectal cancer cell lines were labeled with ¹¹¹chromium and placed into 96-well V-bottomed plates for use as target cells. Patient sera were added at 1:20, 1:100, and 1:1500 dilutions and further incubated for 1 hour. PBMCs were then added to the wells to achieve a 100:1 effector/target ratio. ¹¹¹Chromium released into the supernatant was analyzed by MicroBeta Plus Scintillation counter (Wallac). Cytotoxicity was calculated as percentage of target cell lysis = 100 × (cpm of experimental release – cpm of spontaneous release)/(cpm of maximum release – cpm of spontaneous release).

Analysis of clinical activity. Clinical activity was assessed by applying the RECIST criteria to CT or MRI scans obtained before and after all immunizations. Peripheral blood tumor CEA level was assessed if available. Patients were followed long-term for survival.

Statistics. A positive immune response by ELISpot was defined as described at the 2002 Society of Biologic Therapy Workshop on “Immunologic Monitoring of Cancer Vaccine Therapy”: a T cell response is considered positive if the mean number of spots in 6 wells with antigen exceeds the mean number of spots in 6 control wells by 10 and the difference between the mean of the 6 wells containing antigen and the 6 control wells is statistically significant at a level of P < 0.05 using the Student’s t test (30). A positive response in the intracellular cytokine assay was defined as the percentage of positive cells by FACS analysis after vaccination greater or equal to twice the prevaccination sample. A positive result in the ELISA was defined as a 1:25 antibody titer. Titer is defined as twice the absorbance of the negative control serum.

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