Combining Human and Rat Sequences in Her-2 DNA Vaccines Blunts Immune Tolerance and Drives Antitumor Immunity

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Abstract

Immune tolerance to tumor-associated self-antigens poses a major challenge in the ability to mount an effective cancer vaccine response. To overcome immune tolerance to HER-2, we formulated DNA vaccines that express both human HER-2 and heterologous rat Neu sequences in separate plasmids or as single hybrid constructs that encode HER-2/Neu fusion proteins. Candidate vaccines were tested in Her-2 transgenic (Tg) mice of BALB/c (BALB), BALB/c × C57BL/6 F1 (F1), or C57BL/6 (B6) background, which exhibit decreasing immune responsiveness to HER-2. Analysis of various cocktails or hybrid vaccines defined a requirement for particular combination of HER-2/Neu sequences to effectively prime immune effector cells in HER-2 Tg mice. In B6 HER-2 Tg mice, rejection of HER-2–positive tumors protected mice from HER-2–negative tumors, providing evidence of epitope spreading. Our findings show that a strategy of combining heterologous antigen with self-antigens could produce a potent DNA vaccine that may be applicable to other tumor-associated antigens. Cancer Res; 70(1); 119–28. ©2010 AACR.

Introduction

Dysregulated signal transduction from overexpressed or mutated ErbB-2 (Her-2/neu) leads to cellular immortalization, neoplastic transformation, and tumor progression (1, 2). Consequently, Her-2 is a negative prognostic factor in breast cancer (3–6). Monoclonal antibodies (mAb) such as trastuzumab (Herceptin) and receptor tyrosine kinase inhibitors such as lapatinib are effective against Her-2–positive breast cancers (7–9), but patients typically develop resistance after treatment, suggesting compensatory activity through alternative signaling pathways (10–13).

Her-2 DNA vaccines have shown promising activities in human Her-2 or rat neu transgenic (Tg) mice, but immune tolerance to tumor-associated self-antigens continues to be a major challenge (14, 15). In several experimental autoimmune disease models, tolerance to self-antigens is overcome by immunization with heterologous (xenogeneic) antigens, e.g., murine experimental autoimmune encephalomyelitis is induced by bovine or guinea pig myelin basic protein (16, 17), collagen-induced arthritis by bovine, porcine, or human collagen (18), and autoimmune myasthenia gravis by Torpedo californica acetylcholine receptor (19, 20). These heterologous antigens may share common or cross-reactive epitopes with self-antigens to overcome immune tolerance and trigger self-reactive T cells. Importantly, human or mouse tyrosinase DNA prolongs the survival of dogs with malignant melanoma (21), resulting in the licensing of a DNA vaccine for canine melanoma (21, 22).

To test if heterologous antigens could be exploited to improve Her-2 vaccine efficacy, BALB NeuT mice expressing a rat neu transgene were electrovaccinated with human Her-2 or self neu DNA (23). Greater T-cell response to neu was induced by heterologous Her-2 than by self neu. But only self neu, not heterologous Her-2 vaccine, induced the anti-neu antibodies that were critical in delaying spontaneous tumor formation. These results reflect the exquisite specificity of anti-neu antibodies and that humoral immunity is the primary effector mechanism against neu-induced spontaneous tumors. Not all Her-2–positive breast cancers are sensitive to antibody-mediated destruction (24). We have shown that TUBO and Bam1a cells, established from BALB NeuT spontaneous mammary tumors, were highly sensitive to anti-neu mAb or receptor tyrosine kinase inhibitor. On the other hand, receptor tyrosine kinase inhibitor–resistant BamIR-5 or neu-transfected D2F2/neu cell lines were refractory to antibody therapy. Nevertheless, all test cells could be controlled by CTL, demonstrating the importance of inducing both T-cell and B-cell responses with Her-2 vaccines (24). In this study, we tested the hypothesis that cocktail or hybrid vaccines containing both self human Her-2 and heterologous rat neu DNA sequences would induce greater antitumor immunity than either single antigen.

We have shown that human Her-2 Tg mice on a BALB/c background were much more responsive to Her-2 vaccination
than those on a C57BL/6 background (15). Therefore, the candidate vaccines were tested in Her-2 Tg mice of BALB/c, C57BL/6, or (BALB/c × C57BL/6) F1 background to represent patients with heterogeneous genetic makeup.

Materials and Methods

Mice. All animal procedures were conducted in accordance with accredited institution guidelines and the U.S. Public Health Service Policy on Humane Care and Use of Laboratory Animals. C57BL/6 (B6) and BALB/c (BALB) female mice were purchased from Charles River Laboratory. Heterozygous C57BL/6 Her-2 Tg mice (B6 Her-2 Tg), which expressed the full-length, wild-type human Her-2 under the whey acidic protein promoter were generated in our laboratory and have been previously described (23). pEFBos/granulocyte-macrophage colony-stimulating factor (pGM-CSF) encoding murine GM-CSF was provided by Dr. N. Nishisaki at Osaka University, Osaka, Japan. The blank vector, pVax1 (Invitrogen), was the control.

DNA vaccine constructs. pVax/E2TM (pE2TM) and pcDNA/NeuTM (pNeuTM) encoding the extracellular and transmembrane domains of human (Her-2) or rat (neu) erbB-2, respectively, have been previously described (23). pEFBos/granulocyte-macrophage colony-stimulating factor (pGM-CSF) encoding murine GM-CSF was provided by Dr. N. Nishisaki at Osaka University, Osaka, Japan. The blank vector, pVax1 (Invitrogen), was the control.

Hybrid DNA vaccines pE2-NeuTM (alternatively designated as HuRT) in EP08004513.1 and manuscript submitted by Quaglino and colleagues, pNeuE2TM (alternatively designated as RHuT in USP200702990023 and manuscript submitted by Quaglino and colleagues), and pcDNA/NeuTM were generated (Fig. 1A), using strategies described in Supplementary Fig. S1.

Characterization of DNA constructs. DNA constructs were transiently transfected into 3T3 cells using LipofectAMINE 2000 (Invitrogen) following the instructions of the manufacturer. The mAb TA-1 (Ab5; Calbiochem), N12, N29 (both hybridomas provided by Dr. Yosef Yarden, Weissman Institute; refs. 26, 27), and trastuzumab (Genentech; ref. 28) were used to characterize Her-2 epitopes in the chimera fusion protein refs. 26, 27), and trastuzumab (Genentech; ref. 28) were used to characterize Her-2 epitopes in the chimera fusion protein. The mAb TA-1 (Ab5; Calbiochem), N12, N29 (both hybridomas provided by Dr. Yosef Yarden, Weissman Institute; refs. 26, 27), and trastuzumab (Genentech; ref. 28) were used to characterize Her-2 epitopes in the chimera fusion protein. The mAb TA-1 (Ab5; Calbiochem), N12, N29 (both hybridomas provided by Dr. Yosef Yarden, Weissman Institute; refs. 26, 27), and trastuzumab (Genentech; ref. 28) were used to characterize Her-2 epitopes in the chimera fusion protein.

Cell lines. The BALB/c mouse mammary tumor line D2F2 was established in our group from a spontaneous tumor that arose from the prolactin-induced hyperplastic alveolar nodule line, D2, as previously described (30). D2F2/E2 was generated by stable transfection of D2F2 cells with wild-type human Her-2 (23, 31). Their mouse origin was verified by spectral karyotyping in 2008 (24). E0771/E2 (25) was the control.

C57BL/6 mammary tumor E0771 with wild-type Her-2 (15). E0771 and E0771/E2 developed into palpable tumors in mice of C57BL/6 origin. BALB/c antigen-presenting cells 3T3/KB expressed H2Kd and B7.1 (CD80), and 3T3/EKB or 3T3/NKB expressed additional Her-2 or neu (23, 32). C57BL/6 TC-1 cells (Dr. T.C. Wu, The Johns Hopkins University, Baltimore, MD) expressed endogenous H2Kd and B7.1 (15). TC-1/E2 was transfected to express wild-type Her-2 as antigen-presenting cells. SKOV3, a human ovarian cancer cell line with amplified Her-2 (33), was purchased from American Type Culture Collection and Her-2 expression was verified regularly by flow cytometry. The hybridoma line PC61 (American Type Culture Collection), which produces rat mAb to mouse CD25 was propagated in severe combined immunodeficient mice. All cell lines were maintained as previously described (34).

Immunization. Mice received DNA electrovaccination as we previously described (34). Wild-type mice were immunized twice whereas Her-2 Tg mice were immunized four times. To deplete CD25+ regulatory T cells (Treg), indicated groups of mice received 0.5 mg of CD25 mAb, PC61, Ip, 7 d prior to the first electrovaccination. Depletion of CD25+ T-cells was verified by flow cytometry.

Measurement of Her-2 or neu binding antibodies by flow cytometry. Anti–Her-2 antibodies were measured by binding to Her-2 overexpressing SKOV3 cells using flow cytometry, and antibody concentrations were calculated by regression analysis using mAb TA-1 as the standard (35). Normal mouse serum or isotype-matched mAb was the negative control. Anti-neu antibody was measured with 3T3/NKB cells (Jackson ImmunoResearch) and the standard curve was generated using mAb 7.16.4 (Ab4). Differences in antibody concentration were analyzed by Student’s t test. IgG isotypes were measured with FITC-conjugated goat anti-mouse IgG1 or IgG2a (Jackson ImmunoResearch) and expressed as mean channel fluorescence.

Measurement of T-cell response by IFN-γ ELISPOT. All ELISPOT reagents were purchased from BD Biosciences, unless otherwise specified. Her-2–reactive T cells were enumerated by IFN-γ ELISPOT assay as we described (15, 23). Peripheral blood (PBL) or spleen cells were incubated for 48 h with the antigen-presenting cells TC-1, TC-1/E2, 3T3/KB, or 3T3/EKB at an antigen-presenting cell/lymphocyte ratio of 1:10. The results were expressed as the number of cytokine-producing cells per 10⁶ PBL or spleen cells. Neu-reactive T cells were monitored using 3T3/NKB cells as antigen-presenting cells. Data were analyzed using Student’s t test.

Tumor growth inhibition assay. To measure protective immunity, Her-2 Tg mice were inoculated s.c. in the flank with 2 × 10⁵ D2F2/E2 or E0771/E2 tumor cells 2 wk after the fourth electrovaccination. Tumor growth was monitored weekly, and tumor volume was calculated by XY² / 2 (X = long axis, Y = short axis). To test therapeutic potential, mice were inoculated with 2 × 10⁵ D2F2/E2 or E0771/E2 tumor cells 11 d after a single vaccination. B6 Her-2 Tg mice were depleted of Treg using 0.5 mg of PC61 7 d prior to the first electrovaccination. Starting 3 d following tumor inoculation, mice received three weekly vaccinations. Tumor growth was...
monitored by weekly palpation and difference in tumor incidence was analyzed by the log rank test.

Results

Induction of Her-2 immunity with self versus heterologous DNA vaccine

To compare the activity of self versus heterologous vaccines, BALB NeuT or BALB Her-2 Tg mice were electrovaccinated four times, every 2 weeks, with 50 μg of pE2TM or pNeuTM. In all vaccination experiments, equal amounts of pGM-CSF were coadministered. Antibody response was measured 2 weeks after the final vaccination. As shown in Table 1, pNeuTM induced higher levels of neu binding antibodies when administered as a foreign antigen in Her-2 Tg mice (54 ± 5 μg/mL) rather than as a self-antigen in NeuT Tg mice (16 ± 2 μg/mL), but neither antiserum recognized Her-2, showing antigen specificity. Similarly, pE2TM induced more Her-2–binding antibodies in NeuT (40 ± 3 μg/mL) than in Her-2 Tg (20 ± 5 μg/mL) mice, again neither serum recognized neu. Therefore, antibodies induced by DNA vaccines showed exquisite specificity to the cognate antigen.

In the T-cell compartment, however, heterologous antigen was more effective in activating self-reactive T cells. In NeuT mice, self pNeuTM induced 294 ± 50 spots/10^6 spleen cells of neu-specific, IFN-γ–producing T cells, but heterologous pE2TM activated more anti-neu T cells at 567 ± 176 (P <
0.001). Similarly, in Her-2 Tg mice, heterologous pNeuTM induced 292 ± 16 anti–Her-2 T cells, significantly more than that by self pE2TM (124 ± 4, P < 0.001). Therefore, heterologous Her-2 or neu DNA is superior in overcoming T-cell tolerance. All ELISPOT assays were performed with pooled PBL except in NeuT mice whose splenocytes were used because neu-reactive T cells were not detectable in their PBL.

### Induction of Her-2 immunity with a cocktail vaccine of pE2TM and pNeuTM

The disparate cellular and humoral immunity induced by heterologous and self-antigens led to the hypothesis that antitumor immunity may be enhanced by combined vaccination with rat neu and human Her-2. To test this hypothesis, BALB Her-2 Tg mice were immunized four times with a cocktail of 25 μg of pE2TM and 25 μg of pNeuTM (Fig. 1). Control groups received 50 μg of either pNeuTM or pE2TM. The cocktail vaccine induced 74 ± 9 μg/mL of Her-2 binding antibodies at week 8, compared with 23 ± 17 μg/mL by pE2TM alone, showing significant amplification of humoral response by the presence of heterologous neu in the vaccine (Fig. 1B). In the T-cell compartment, pE2TM activated more T cells than heterologous pNeuTM after two vaccinations (Fig. 1C). After four vaccinations, T-cell response in pNeuTM immunized mice escalated to reach 292 ± 16/10^6 PBL, whereas the pE2TM immunized mice produced 116 ± 21/10^6 PBL. Surprisingly, T-cell response in the cocktail vaccine group was only comparable to that in pE2TM group, below that induced by the heterologous pNeuTM. Because pNeuTM itself did not induce Her-2–binding antibodies (Table 1; ref. 23) and Her-2–specific T-cell response was not elevated by the cocktail vaccine (Fig. 1C), the enhanced Her-2 antibody response could be due to immune stimulation by the foreign epitopes in neu.

All test and control mice were challenged s.c. with D2F2/E2 tumor cells (Fig. 1D), which were rejected in 72% (five of seven) of mice receiving the pE2TM + pNeuTM cocktail, compared with 38% (three of eight) in the pE2TM group (P < 0.05) and 60% (6 of 10) in the pNeuTM group. The elevated levels of Her-2 antibodies induced by the cocktail vaccine might mediate antibody-dependent cell-mediated cytotoxicity to exert greater inhibition of tumor growth even though D2F2/E2 tumor cells are insensitive to the direct inhibitory effects of Her-2 antibodies (24). Importantly, pNeuTM, which induced strong T-cell response without anti–Her-2 antibody, showed comparable antitumor activity to the cocktail vaccine.

### Generation and characterization of hybrid Her-2-neu vaccines

Because antibody-sensitive and -resistant cells both exist in human breast cancer, the efficacy of combining Her-2 and neu into one vaccine formulation was further tested with hybrid vaccines that encode fusion proteins of Her-2 and neu (Fig. 1D; Supplementary Fig. S1). The chimeric E2-NeuTM protein is a transmembrane Her-2-neu fusion protein with a truncated ICD. Neu-E2TM is the reverse product with neu at the NH2 terminus and Her-2 at the COOH terminus (36). The third chimeric vaccine pE2-Neu500TM was modified from pE2-NeuTM to restore Her-2 amino acids 501 to 687 which encompassed the epitope(s) recognized by trastuzumab (28). The fusion protein generated from each test construct was verified after transient transfection into 3T3 cells, using a panel of Her-2 and neu mAbs as well as polyclonal antibodies (Supplementary Fig. S1).

Immunogenicity of the hybrid vaccines was verified in wild-type BALB/c females. Comparable levels of Her-2 antibodies (40 μg/mL) were induced by pE2TM, pE2-NeuTM, and pE2-Neu500TM (Supplementary Fig. S2A), each containing Her-2 NH2-terminal codons. pNeu-E2TM containing NH2-terminal neu induced less Her-2 binding antibodies at 18 ± 4 μg/mL; pNeuTM did not induce Her-2 binding antibodies as predicted (Table 1). Conversely, ~70 μg/mL of neu-binding antibodies were induced by pNeuTM or pE2-NeuTM, which contained neu in the NH2 terminus (Supplementary Fig. S2A) with low-level activity from pE2-Neu or pE2-Neu500TM (11 ± 3 and 6.3 ± 3 μg/mL, respectively). Therefore, in wild-type mice, fusion proteins which contained Her-2 or neu in the NH2 terminus induced greater antibody response to the respective antigen.

Regarding T cells, high levels of Her-2–specific IFN-γ–secreting cells were induced by pE2TM or pE2-NeuTM and modestly by pNeuTM or pNeu-E2TM (Supplementary Fig. S2B). Neu-specific T-cell response was comparable in mice vaccinated with pNeuTM, pNeu-E2TM, or pE2TM, with somewhat weaker response from pE2-NeuTM. Bearing some variation, hybrid vaccines were immunogenic in wild-type mice,

### Table 1. Induction of Her-2 or neu immunity with self vs. heterologous DNA vaccination

<table>
<thead>
<tr>
<th>antibodies (μg/mL)</th>
<th>pNeuTM vaccinated</th>
<th>pE2TM vaccinated</th>
</tr>
</thead>
<tbody>
<tr>
<td>BALB NeuT mice</td>
<td>BALB Her-2 Tg mice</td>
<td>BALB NeuT mice</td>
</tr>
<tr>
<td>Anti-Neu antibodies</td>
<td>16 ± 2*</td>
<td>0.3 ± 0.2</td>
</tr>
<tr>
<td>Anti-Her-2 antibodies</td>
<td>0*</td>
<td>20 ± 5</td>
</tr>
<tr>
<td>IFN-γ T cells (spots/10^6 cells)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neu antigen</td>
<td>294 ± 50†</td>
<td>567 ± 176</td>
</tr>
<tr>
<td>Her-2 antigen</td>
<td>22 ± 4</td>
<td>1,014 ± 128</td>
</tr>
</tbody>
</table>

*P < 0.001 compared with the same strain of mice receiving pE2TM.
†P < 0.05 compared with the same strain of mice receiving pE2TM.
with the NH₂ terminus sequences of Her-2 showing greater immunogenicity.

Activity of hybrid vaccines in Her-2 Tg mice

**BALB Her-2 Tg mice.** The immunogenicity and efficacy of hybrid vaccines were tested in BALB Her-2 Tg mice using pE2-NeuTM or the reverse pNeu-E2TM. pE2TM was the control. pE2-NeuTM induced significantly higher levels of Her-2 antibodies, compared with pE2TM or pNeu-E2TM, more prominently after the third vaccination (Fig. 2A), consistent with greater immunogenicity of epitopes in Her-2 NH₂ terminus. Immunoglobulin isotype analysis showed the induction of both IgG₁ and IgG₂a (Fig. 2A, right).

Her-2-reactive T cells in pooled PBL were measured after two and four vaccinations (Fig. 2B). After two vaccinations, pE2TM induced more IFN-γ-secreting T cells than either hybrid constructs. After four vaccinations with pE2-NeuTM, however, hybrid pE2-NeuTM showed a modest edge over pE2TM (153 ± 4/10⁶ compared with 116 ± 21/10⁶) and pNeu-E2TM (92 ± 4/10⁶), suggesting the advantage of concurrent presentation of neu and preservation of Her-2 NH₂ terminus in the vaccine regimen.

When mice were challenged subcutaneously with D2F2/E2 cells, 80% (8 of 10) of pE2-NeuTM–vaccinated mice were protected, but only 40% of pE2TM–vaccinated mice (2 of 5) and 33% (3 of 9) of pNeu-E2TM–vaccinated mice were protected (Fig. 2C). Thus, enhanced Her-2 antibody and T-cell response was induced after four vaccinations with hybrid pE2-NeuTM to result in elevated antitumor activity.

**(BALB × B6) F1 Her-2 Tg mice**

We previously reported that BALB Her-2 Tg mice were much more responsive to pE2TM than B6 Her-2 Tg mice, and that (BALB × B6) F1 Her-2 Tg mice produced intermediate response (15, 25). To further test the efficacy of hybrid vaccines, F1 Her-2 Tg mice which showed intermediate Her-2 reactivity were immunized with pE2TM, pE2-NeuTM, or pE2-Neu500TM; the latter was constructed to restore Her-2 amino acids 501 to 650, thus reintroducing the epitopes recognized by trastuzumab and reducing the neu sequence from 263 amino acids to 110 amino acids. In some test mice, Treg were depleted with CD25 mAb 1 wk before vaccination (Fig. 3A; ref. 15).

In mice with intact Tregs, Her-2 antibodies induced by pE2-NeuTM more than doubled that by pE2TM or pE2-Neu500TM (Fig. 3B, left). After Treg depletion, antibody response increased most significantly in the pE2-Neu500TM group to approach that by pE2-NeuTM (Fig. 3B, right). These

![Figure 2](https://example.com/figure2.png)

**Figure 2.** Hybrid vaccines in BALB Her-2 Tg mice. A, mice were vaccinated four times with pE2TM, pE2-NeuTM, or pNeu-E2TM. Her-2 antibodies in individual mice were measured after each vaccination. IgG₁ and IgG₂a levels were analyzed after the last vaccination. B, Her-2-reactive IFN-γ-secreting cells in pooled PBL were measured after the second and fourth vaccination. C, at 2 wk following the last vaccination, mice were challenged with 2 × 10⁵ D2F2/E2 and tumor growth was monitored by weekly palpation. There were 4 to 10 mice per group (*, P < 0.001).
results suggested that Her-2 extracellular domain residues 500 to 687 may activate Treg and their presence in pE2-Neu500TM might compromise vaccine activity.

After four vaccinations, pE2-NeuTM induced significantly more IFN-γ–producing T cells than pE2TM or pE2-Neu500TM (Fig. 3C, left), consistent with Treg activation by pE2-Neu500TM. Furthermore, the shorter 110 residues of heterologous neu in pE2-Neu500TM may be insufficient to overcome Her-2 tolerance. After Treg depletion, T-cell response in every group increased; still, pE2-NeuTM induced more than twice the number of IFN-γ–producing T cells (397 ± 30) than pE2TM (147 ± 15). T-cell response to pE2-Neu500TM improved, but did not approach pE2-NeuTM, indicating the need for additional neu residues for Her-2 T-cell activation (Fig. 3C).

To measure antitumor immunity, the experiment was repeated with an increased number of mice. Treg-depleted F1
Her-2 Tg mice were vaccinated four times with pE2-NeuTM, pE2TM, or control pVax1 before they were inoculated s.c. with Her-2–transfected D2F2/E2 cells of BALB/c origin or E0771/E2 cells of C57BL/6 origin. There were six to seven mice in each group. All tumors grew progressively in pVax1 control mice (Fig. 3D). pE2TM vaccination protected six of seven mice from D2F2/E2 and five of seven mice from E0771/E2. In the pE2-NeuTM–vaccinated group, all mice rejected the challenging tumor, either D2F2/E2 or E0771/E2. Therefore, the hybrid pE2-NeuTM induced stronger humoral and cellular immunity than pE2TM in F1 Her-2 Tg mice to provide complete protection from tumors of either parental background.

B6 Her-2 Tg mice

pE2-NeuTM was further tested in B6 Her-2 Tg mice which were poorly responsive to pE2TM vaccination (15, 23). Treg were depleted with CD25 mAb 1 week before mice were vaccinated with pE2-NeuTM or control pE2TM.

After a single vaccination, pE2-NeuTM already induced significantly more Her-2–binding antibodies than pE2TM. After four vaccinations, pE2-NeuTM induced 64 ± 4 μg/mL of Her-2 antibodies and pE2TM induced 20 ± 9 μg/mL (Fig. 4A). In T-cell response, pE2TM induced 13 ± 2 IFN-γ–secreting cells/106 PBL, whereas the pE2-NeuTM hybrid induced 49 ± 4 cells (Fig. 4B). At 2 weeks after the final vaccination, mice were challenged with 2 × 10^5 E0771/E2 cells. In 40% (two of five) of pE2TM–vaccinated mice and 100% (five of five) of pE2-NeuTM–vaccinated mice, the tumors were rejected, further supporting greater efficacy of the hybrid vaccine (Fig. 4C, left). To determine whether rejection of E0771/E2 resulted in immune priming to other tumor-associated antigens, pE2-NeuTM–vaccinated mice which rejected the primary tumors were re-challenged 7 weeks later with parental E0771 which did not express Her-2 and 80% (four of five) of these mice rejected the second tumor challenge (Fig. 4C, right). Thus, the pE2-NeuTM hybrid overcame Her-2 immune tolerance regardless of their intrinsic level of tolerance to Her-2, inducing tumor rejection while triggering immune priming to additional tumor-associated antigens.

Efficacy of one preventive and three therapeutic vaccinations

To further test the efficacy of pE2TM and pE2-NeuTM in more stringent conditions, BALB Her-2 and B6 Her-2 Tg mice were electrovaccinated once before they were inoculated s.c. with D2F2/E2 and E0771/E2 cells, respectively (Fig. 5A). Mice then received three weekly vaccinations following tumor inoculation. In B6 Her-2 Tg mice, Treg were depleted before the first vaccination. Her-2 antibody production was evident after a single vaccination with the hybrid DNA and tumor inoculation. Antibody levels continued to escalate with repeated vaccination (Fig. 5B and D, left). In both strains of mice,
pE2-NeuTM showed significantly greater immunogenicity than pE2TM. In BALB Her-2 Tg mice, tumor growth was inhibited by 88% (pE2-NeuTM) and 70% (pE2TM), showing that one preventive and three therapeutic vaccines provided significant antitumor immunity. Primary cultures of D2F2/E2 tumor outgrowths were prepared from each test group and Her-2 expression was retained by the outgrowth, whether or not mice were vaccinated.

In B6 Her-2 Tg mice, however, significant rejection of E0771/E2 tumors was not achieved by either vaccine (3 of Figure 5. The efficacy of one preventive and three therapeutic vaccines. A, BALB Her-2 Tg mice received $2 \times 10^5$ D2F2/E2 tumors 11 d after a single protective vaccination. Mice then received three weekly vaccinations, starting 3 d after tumor inoculation. B, Her-2 antibodies were measured after each vaccination and tumor growth was monitored weekly. There were six to eight mice per group. C, expression of Her-2 by primary cultures of D2F2/E2 tumor was measured by mAb TA-1. D, B6 Her-2 Tg mice were inoculated with $2 \times 10^5$ E0771/E2 tumor cells following the same vaccination scheme, except that CD25 mAb was given 1 wk before the first vaccination. Inset, Her-2 expression on E0771/E2 tumor outgrowth in pVax1 group. There were 9 to 10 mice per group (*, $P < 0.05$; **, $P < 0.01$).
10 in pE2TM-vaccinated and 1 of 10 in pE2-NeuTM-vaccinated mice) although high levels of Her-2 antibodies were induced by pE2-NeuTM. However, the average volume of tumors in immunized mice was significantly reduced compared with the pVax1 group. When primary cultures of E0771/E2 tumors were prepared at week 7, little to no Her-2 expression was detected even in mice which received the pVax1 control vector. Therefore, loss of Her-2 expression in vivo muted the excellent anti–Her-2 immunity induced by vaccination.

These results showed the efficacy of DNA vaccination against Her-2–expressing tumors when administered as one preventive and three therapeutic vaccines. The vaccine specificity was further illustrated by the rejection of D2F2/E2 tumors which retained Her-2 expression in vivo, but less so of E0771/E2 tumors, which lost Her-2 expression in vivo.

**Discussion**

A new vaccination regimen that presented heterologous rat neu together with self Her-2 in a cocktail or as a hybrid construct significantly improved anti–Her-2 immunity and protective activity in Her-2 Tg mice of three different genetic backgrounds. Most notably, seven of eight tumors in BALB Her-2 Tg mice were rejected after one preventive and three therapeutic vaccination with pE2-NeuTM. Antibody response was highly elevated by the cocktail or pE2-NeuTM hybrid vaccine, even though neu itself did not induce Her-2–binding antibodies, suggesting immune stimulation by the foreign epitopes in neu. On the contrary, heterologous pNeuTM induced significantly more Her-2–reactive T cells than self pE2TM, but elevated T-cell response was evident only after repeated pNeuTM immunization, showing a delayed yet eventually more profound T-cell activation by cross-reactive T-cell epitopes from neu. This disparity in antibody and T-cell responses induced by heterologous Her-2/neu antigen was consistent with our previous report (23).

The NH2 terminus of Her-2 showed greater immunogenicity in both wild-type BALB/c and BALB Her-2 Tg mice, whereas heterologous neu residues 501 to 688 were important in overcoming Her-2 tolerance as both pNeu-E2TM or pE2-Neu400TM lacking neu amino acids 501 to 688 were less effective. After Treg depletion, Her-2 antibody response to pE2-Neu400TM approached that of pNeu-E2TM and may indicate activation of Treg by Her-2 residues 501 to 687. Based on these findings, it may be possible to design hybrid vaccines targeting other tumor-associated antigens once the critical regions for immune activation in self and heterologous antigens are defined.

The high level of sequence identity and the similarity in three-dimensional structures of Her-2 and neu enabled the generation of stable fusion proteins E2-NeuTM, E2-Neu400TM, and Neu-E2TM to facilitate vaccine development. Incorporation of strong foreign epitopes such as tetanus toxin, rather than heterologous neu may also enhance the immune response to Her-2, but without the benefit of shared epitopes. When extrapolating from these findings to new vaccines, the high level of sequence identity between Her-2 and neu (88%) or between Her-2 and mouse ErbB-2 (84%) should be noted. It is possible that only a highly homologous sequence from a xenogeneic origin is effective in overcoming self-tolerance.

IFN-γ–producing T cells can be detected in the PBL of Her-2 Tg mice. It is therefore possible to measure T-cell activity with pooled PBL and monitor tumor growth in the same group of mice. In NeuT mice, IFN-γ–producing T-cells were not found in the PBL although they were readily detected in the spleens (Table 1). It is possible that circulating neu-specific T cells extravagated and localized in the 10 mammary glands which expressed large quantities of neu, and thus, are below detection in PBL.

When pE2-NeuTM was administered as one preventive followed by three therapeutic vaccinations, D2F2/E2 tumors which retained Her-2 expression in vivo were rejected in seven of eight BALB Her-2 Tg mice, demonstrating the efficacy of DNA vaccine in this more stringent setting. Comparable levels of Her-2 antibodies were induced in B6 Her-2 Tg mice following the same vaccination regimen after Treg depletion. Although average tumor growth was slowed in pE2-NeuTM–immunized mice for several weeks, rejection of the E0771/E2 tumor was poor because these tumor cells lost Her-2 expression in vivo.

Unlike syngeneic mice, humans are heterogeneous in their genetic background. With Her-2 Tg mice in BALB, B6, and F1 background, variation in immune responses to Her-2 vaccine became evident (15). Using hybrid pE2-NeuTM, even the most tolerant B6 Her-2 Tg mice responded with a striking complete protection from tumor challenge. With one preventive and three therapeutic vaccinations, hybrid pE2-NeuTM induced significant tumor rejection in BALB Her-2 Tg mice. The concurrent presentation of self and heterologous antigen in a cocktail or in a hybrid construct augmented anti–Her-2 immunity, and thus, should be considered for future vaccine design.

**Disclosure of Potential Conflicts of Interest**

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References


Correction: Combining Human and Rat Sequences in Her-2 DNA Vaccines Blunts Immune Tolerance and Drives Antitumor Immunity

An early version of this article (Cancer Res 2010;70:119–28) under the title “Human Her-2 DNA Vaccines Containing Heterologous Rat Neu Sequence Overcome Immune Tolerance to Induce Elevated Antitumor Immunity” was published by mistake in the January 1, 2010 issue of Cancer Research (1). The correct version of the article appears online.

Reference


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Combining Human and Rat Sequences in Her-2 DNA Vaccines Blunts Immune Tolerance and Drives Antitumor Immunity

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