Classifying Patients for Breast Cancer by Detection of Autoantibodies against a Panel of Conformation-Carrying Antigens

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Abstract

Patients with breast cancer elicit an autoantibody response against cancer proteins, which reflects and amplifies the cellular changes associated with tumorigenesis. Detection of autoantibodies in plasma may provide a minimally invasive mechanism for early detection of breast cancer. To identify cancer proteins that elicit a humoral response, we generated a cDNA library enriched for breast cancer genes that encode membrane and secreted proteins, which are more likely to induce an antibody response compared with intracellular proteins. To generate conformation-carrying antigens that are efficiently recognized by patients' antibodies, a eukaryotic expression strategy was established. Plasma from 200 patients with breast cancer and 200 age-matched healthy controls were measured for autoantibody activity against 20 different antigens designed to have conformational epitopes using ELISA. A conditional logistic regression model was used to select a combination of autoantibody responses against the 20 different antigens to classify patients with breast cancer from healthy controls. The best combination included ANGPTL4, DKK1, GAL1, MUC1, GFRA1, GRN, and LRRC15; however, autoantibody responses against GFRA1, GRN, and LRRC15 were inversely correlated with breast cancer. When the autoantibody responses against the 7 antigens were added to the base model, including age, BMI, race and current smoking status, the assay had the following diagnostic capabilities: c-stat (95% CI), 0.82 (0.78–0.86); sensitivity, 73%; specificity, 76%; and positive likelihood ratio (95% CI), 3.04 (2.34–3.94). The model was calibrated across risk deciles (Hosmer–Lemeshow, P = 0.13) and performed well in specific subtypes of breast cancer including estrogen receptor positive, HER-2 positive, invasive, in situ and tumor sizes >1 cm. Cancer Prev Res; 7(5); 545–55. ©2014 AACR.

Introduction

For patients with breast cancer, early and personalized diagnosis is crucial for optimizing treatments leading to long-term survival. Although mammography is the most widely used method to detect breast cancer, approximately 20% of screening mammograms result in a false negative diagnosis largely because of high breast density (1). In addition, 1 in 10 women who get a mammogram will need additional imaging (2). Yet, the overwhelming majority of these women will not have breast cancer, as only 2 to 4 of every 1,000 screening mammograms leads to a cancer diagnosis (3). Therefore, there is an urgent clinical need to develop a novel, minimally invasive diagnostic strategy for the early diagnosis of breast cancer.

Measuring the levels of tumor markers, which are materials of genetic origin produced by tumors themselves or by the host in response to the tumor (4, 5), is a promising strategy for the early diagnosis of cancer. At present, there is no established tumor marker that is secreted into the peripheral circulation that can be measured by a blood test for the diagnosis of breast cancer. Currently, tumor markers that are accepted in clinical practice are tissue-based prognostic markers, such as the estrogen receptor (ER), HER-2 amplification, 21-gene Oncotype DX, and 70-gene Mammaprint (6–12). All require an invasive biopsy or surgical procedure to acquire tumor tissue for assessment, bearing a heavy burden on patients. Serum tumor markers are valuable tools that allow minimally invasive procedures for sampling to promote the early diagnosis of cancer as well as following the prognosis after treatment (4, 5). However, tumor markers produced by tumor cells usually have relatively low concentrations in the peripheral circulation, especially in early-stage disease. It has been previously shown that tumor-associated antigens (TAA) can elicit an antibody response in patients with cancer (13–15). For patients with breast cancer, several TAs, including p53, HER-2, MUC1, HSP-60, NY-ESO-1, and c-myc, have been
identified (for review, see refs. 16–18). Because the immune system can produce a considerable amount of antibody even when it is exposed to a limited amount of tumor antigen (19), the detection of antibodies against tumor proteins can be more sensitive than screening for the tumor antigens. However, one of the largest barriers to utilizing anti-TAA antibodies as diagnostic markers is the identification of the tumor antigens recognized by the autoantibodies. Previously, serologic identification of antigens by recombinant expression cloning (SEREX) and phage display methods have been used to identify tumor antigens that elicit an autoantibody response in patients. For both methods, cDNA expression libraries were derived from cancer tissue or cell lines (20–22) and then clones encoding antigens reactive with antibodies in patients’ sera were selected. Because candidate antigens are produced as denatured fragments in bacteria, the antigens lack conformational structures that represent the majority of immunogenic moieties of proteins (23–25). Membrane and secreted proteins require interactions with membrane lipids and/or posttranslational modifications, such as disulfide bond formation, for proper folding. Estimations based on available antigen–antibody complex crystal structures indicate that more than 90% of epitopes on a protein are conformational or discontinuous epitopes that form by spatial proximity (24, 25). Discontinuous epitopes consist of amino acid segments that are distantly separated in the antigen sequence and are brought into proximity by the folding of the protein. Consistent with these limitations, many of the previously identified antigens by phage display methods are nonbiologic peptides derived from noncoding sequences with questionable utility (15). Other proteomic methods for antibody detection have been developed, such as protein microarrays, reverse-capture microarrays, serological protome analysis and nucleic acid programmable protein array (16, 26–28).

Here we report the use of a molecular approach to identify tumor antigen candidates that elicit an antibody response in patients with breast cancer. Previously, we generated a breast cancer cDNA library from membrane-associated polyribosomal (MAP) RNA, which encodes secreted and membrane proteins, and subtracted the library with RNA from normal tissues (29). Secreted proteins are more easily delivered from tumor cells to lymph nodes, where interactions of immune cells take place resulting in abundant high-affinity antibodies. Membrane surface proteins are commonly released in a soluble form from tumor cells through metalloproteinase-dependent cleavage. The shed proteins are more easily transferred to the lymph nodes than intra-cellular proteins (30, 31). Consequently, the obtained subtracted library, referred to as the membrane-associated polyribosomal cDNA library (MAPcL), is enriched with clones encoding membrane and secreted TAA that are highly abundant in breast cancer and should preferentially induce an antibody response in patients (29). In addition, we have established a method for producing recombinant antigens as Fc fusion proteins designed to have native conformations, which is essential for the expression of membrane and secreted proteins that may induce an antibody response in patients.

We have developed a conformation-carrying antigen ELISA-based strategy to discriminate between breast cancer and healthy patients by the detection of autoantibodies against a panel of TAAs. Twenty antigens were selected from the most abundant genes represented in the MAPcL and Fc fusion proteins were generated. Blood was collected from 200 newly diagnosed patients with breast cancer and 200 healthy women as age-matched controls. The 400 plasma samples were screened for the presence of autoantibodies against the 20 different MAPcL-derived antigens using ELISA. A combination of 7 antigens with patient demographics yielded the best positive likelihood ratio to discriminate between healthy and patients with breast cancer.

Materials and Methods

Plasmid construction

For production of MAPcL-rabbit Fc-tagged antigens, 2 constructs, pSecTag2 (Invitrogen) and pFUSE-rIgG-Fc1 (InvivoGen), were both utilized to generate the 20 MAPcL-rFc expression constructs because of restriction site availability for cloning. pSecTag2 was modified by amplifying the Fc portion of rabbit immunoglobulin G (IgG) using primers 5’-CCCGATATCATCGACCCACGTGGC-CACCC-3’ and 5’-AAAGGAAAAAGCGCGCGCCCT-AATTA-CGCCGAGAGCGGGAG-3’ (Integrated DNA Technologies) using pFUSE-rIgG-Fc1 as a template. The rFc PCR product was digested with EcoRV and NotI and inserted into pSecTag2, referred to as pSecTag2-rFc, which contains an IgK signal sequence for secretion. The pFUSE-rIgG-Fc1 contains an IL-2 signal sequence. To keep the signal sequence consistent between the 2 plasmids, the IgK leader sequence was amplified via PCR using pSecTag2 as a template. The IL-2 leader sequence was then replaced with the IgK signal sequence, creating pFUSE-IgK-rFc.

The accession numbers of the 20 MAPcL genes used as templates for cloning and predicted signal sequences are indicated in Table 1. The signal sequences of each encoded protein were determined using SignalP (32, 33). If a protein contained a transmembrane domain, only the encoded extracellular portion was included. The transmembrane domains were predicted using the TMHMM database (34). The amino acid numbers encoded by the cloned fragment are shown in Table 1. ANGPTL4, CDH3, DKK1, SPON2, SSRI, CST2, GFRA1, and GAL1 were custom cloned into pSecTag2-rFc using the SfiI and KpnI restriction sites (Genscript). EPHA2, IGBPBP2, and LAMC2 were custom cloned into pSecTag2-rFc using the KpnI and BamHI restriction sites. GRN, MUC1, and LRRC15 were custom cloned into pSecTag2-rFc using the SfiI and BamHI restriction sites. HER-2, LRPI0, SPINT2, and SUSD2 were cloned into pFUSE-IgK-rFc using the SfiI and XhoI restriction sites. CD147 was cloned into pFUSE-IgK-rFc using the BamHI and SacI restriction sites. CD320 was cloned into pFUSE-IgK-rFc using the EcoRI and XhoI restriction sites.
For production of His-tagged HER-2, HER-2 was amplified via PCR using primers 5′-CCC
AAGCTTGACCCACCCCAAGTGTGCACCGGCAC-3′ and 5′-GTGCTCGAGTCACGTC-AGAGGGCTGGCTCTCTGCTCG-3′. The product was digested with HindIII and XhoI and cloned directionally into the pET-28a expression vector.

Cell culture
293T and SKBR3 cell lines were cultured in DMEM with 10% FBS. Cultures were maintained at 37°C with 5% CO2 in a humidified incubator. All cell lines were authenticated and tested negatively for mycoplasma.

Protein production
The MAPcL–rFc fusion proteins were produced in 293T cells. Briefly, 293T cells were transfected using Effectene (Qiagen) according to manufacturer’s specifications. During transfection, the cells were cultured in DMEM with 2% FBS. Supernatants containing the secreted fusion proteins were harvested, centrifuged to clear cell debris, and supplemented with 0.1% sodium azide. His-HER-2 was produced in Escherichia coli (E. coli) BL21 (Invitrogen) and purified using IMAC affinity chromatography.

Sandwich ELISA
Microtiter plates (Nalge Nunc) were coated overnight with 2 μg/mL goat anti-rabbit Fc (Jackson Immunoresearch) diluted with phosphate buffered saline. The supernatants containing the rFc fusion proteins were diluted 1:3 serially in standard blocking buffer (0.5% bovine serum albumin and 0.1% sodium azide in PBS). Plates were washed once, and the serially diluted supernatants were transferred to the microtiter plates. Rabbit IgG of known concentration was diluted similarly and added to one row of the microtiter plate in order to quantify the amount of fusion protein present in the culture media. After incubating for 2 hours, plates were washed twice and 50 μL of horse-radish peroxidase (HRP)-conjugated goat anti-rabbit IgG (Jackson Immunoresearch) diluted 1:3,000 in standard blocking buffer with 0.05% Tween 20 added. After a 2-hour incubation, plates were washed 4 times and developed with 100 μL/well of 3,3',5,5'-tetramethylbenzidine (TMB) substrate (Pierce). The development reaction was stopped after 5 minutes with 50 μL/well of 2N H2SO4, and the absorbance was measured at 450 nm to determine the concentration. The absorbance at 690 nm was subtracted to remove background signal.

Antibody recognition of conformational versus denatured HER-2 protein
For the conformational HER-2 assay, microtiter plates were coated with 2 μg/mL goat anti-rabbit Fc (Jackson Immunoresearch) in PBS overnight. HER-2-ECD-rFc was then added to each well, 100 μL/well. For denatured HER-2,
microtiter plates were coated with 2 μg/mL His-HER-2-ECD in PBS overnight.

Three HER-2 antibodies were used in the assay: anti-HER-2 3F27 (US Biological), anti-HER-2 3F32 (US Biological), and Herceptin (Genentech). Each antibody was diluted to 1 μg/mL in standard blocking buffer with 0.05% Tween 20. The antibodies were then serially diluted. After washing once, 50 μL/well of the serially diluted antibodies was added to the plates and incubated for 2 hours at room temperature. The plates were washed 3 times, and species appropriate HRP-conjugated secondary antibodies were added at a 1:3,000 dilution. Plates were washed 4 times and developed with 100 μL/well TMB substrate for 5 minutes. Development was stopped with 50 μL/well 2N H2SO4. Absorbance was measured at 450 nm, and the 690 nm absorbance was subtracted to account for background.

The same antibodies were used to stain HER-2 in SKBR3 breast cancer cells via flow cytometry. SKBR3 cells were detached from dish using Cell Dissociation Solution Non-enzymatic 1 x (Sigma, catalog no. C5914). A total of 2 x 10^5 cells were incubated with 0.5 μg/mL of each antibody for 1 hour at room temperature. The cells were then washed, and a 1:200 dilution of PE-conjugated antibody for the appropriate species was added. The cells were again washed, resuspended in FACS buffer (PBS with 5% bovine serum albumin and 0.1% sodium azide) and analyzed by flow cytometry.

**Competition of Herceptin binding**

Microtiter plates were coated with 4 μg/mL goat anti-rabbit Fc and incubated overnight. After 1 wash, 100 μL/well HER-2-ECD-rFc was added to each well and incubated overnight. HER-2-Fc and CD30-Fc chimeric proteins (R&D Systems) were serially diluted from a starting concentration of 10 μg/mL. Herceptin was added to a final concentration of 10 ng/mL in each of the serial chimeric protein dilutions. Plates were washed twice, and 50 μL/well of chimeric protein/Herceptin mixture was applied to the plate. Plates were then washed 3 times, and a 1:3,000 dilution of HRP goat anti-human IgG was applied to each well, 50 μL/well. After 4 washes, 100 μL/well TMB substrate was added to each well. Development was stopped with 50 μL/well 2N H2SO4 after 5 minutes. Absorbance was measured at 450 nm with 690 nm absorbance subtracted.

**Patients**

The inclusion criteria for cases were women more than 30 years of age that were newly diagnosed with breast cancer (any type) at Sanford Health, Sioux Falls, SD. Patients were asked to provide one extra 10 mL EDTA tube of blood before mastectomy, lumpectomy, radiation therapy, chemotherapy, or other treatment. Case subjects were excluded only if they had a previous history of cancer of any kind. Healthy control subjects had a negative mammogram within 6 months before the blood draw. Healthy subjects were excluded if there was a history of previous cancer of any kind or a history of autoimmune disease. All patients provided written informed consent, and the Sanford Health IRB approved the study protocol. Blood samples from 200 patients with breast cancer were collected from October 8, 2009, to April 17, 2012. In addition, 200 age-matched healthy control blood samples were collected from October 16, 2009, to January 19, 2011. See Table 2 for enrolled patients’ characteristics.

**Serum collection**

Blood was collected in a 10 mL EDTA tube and centrifuged at 2,000 x g for 10 minutes. Plasma was removed from the tube, aliquoted and stored at −80°C until screening for the presence of autoantibodies.
Conformation-carrying antigen ELISA

Microtiter plates (Nalge Nunc) were coated overnight with 4 μg/mL goat anti-rabbit Fc (Jackson Immunoresearch) in PBS. Plates were washed once, and 100 μL/well of MAPcL–rFc fusion protein was added. Plates were incubated for 2 hours and washed twice. The plates were then coated with 50 μL/well of optimized blocking buffer (PBS with 0.5% bovine serum albumin, 0.2% dry milk, 0.1% polyvinylpyrrolidone, 20 mmol/L L-glutamine, 20 mmol/L L-Arginine, 0.1% sodium azide, 10% goat serum, and 0.05% Tween 20). The plates were incubated for 1 hour at 37°C and washed once. Serum samples diluted 1:100 in optimized blocking buffer were added and incubated for 2 hours at room temperature. Plates were then washed 3 times, and autoantibodies were detected using an HRP-conjugated goat anti-human IgG (Jackson Immunoresearch) diluted 1:3,000 in standard blocking buffer with 0.05% Tween 20. Plates were incubated for 1 hour at room temperature, washed 4 times, and developed with 100 μL/well of TMB substrate for 15 minutes. Development was stopped with 50 μL/well 2N H₂SO₄, and the absorbance was measured at 450 nm. The absorbance at 690 nm was subtracted to remove background signal. Each 96-well plate included 14 samples from breast cancer subjects and 14 samples from normal mammogram subjects. Each sample was tested in triplicate within the same plate. One row in each plate was subjected only to blocking buffer as a negative control for the ELISA (Supplementary Fig. S1).

Statistical methods

Controls were individually matched to 200 patients with breast cancer 1:1 within a 3-year age window using a greedy caliper matching algorithm (35) while blinded to assay data. For each subject, the antigen level was transformed by subtracting the mean of the triplicate measurements. If the difference was less than zero, it was set to zero, and the square root was taken to yield a more symmetrical distribution.

Differences in demographics and autoantibody responses between patients breast cancer and controls were tested using 2-sample t test and χ² test for continuous and categorical data, respectively. The incremental improvement to the c-statistic [i.e., concordance index, area under the receiver-operating characteristic (ROC) curve] was tested by adding the autoantibody response to each antigen to a logistic regression model that already included age, BMI, race, and current smoking status. The model calibration was tested using the Hosmer–Lemeshow goodness-of-fit measure, which constructs a χ² statistic by comparing the predicted and observed number of cases by probability decile (36).

After assessing the individual antigens, a multivariable conditional logistic regression analysis with strata for age-matching was used to determine the subset of antigens that minimized Akaike’s Information Criterion (37); all models were adjusted for BMI, race, and current smoking status. Exploratory subgroup analyses were performed to determine if the multivariable subset of antigens performed differently in a particular type of breast cancer. The multivariable model was tested in the following subgroups: invasive, in situ, ER positive, tumor maximum dimension >1 cm, lymph node involvement, and HER-2 positive. The critical level α was set to <0.05/20 antigens = 0.0025 using the Bonferroni correction. SAS version 9.3 software was used for all analyses.

Results

Generation of tumor-associated antigens designed to have native conformations

To identify TAAs that elicit a humoral response in patients, candidate genes that encode membrane and secreted proteins were selected from the most abundant genes represented in the MAPcL. Because only 10% of epitopes on proteins are in a linear continuous sequence (24), we utilized a eukaryotic expression system to generate conformation-carrying tumor antigens that are properly folded and contain noncontinuous epitopes for use in the detection of autoantibodies. Sequences encoding the extracellular domains (ECD) or the secreted proteins without the signal sequence of the candidate MAPcL genes were cloned into the ECD of HER-2 were analyzed: the conformation-carrying HER-2-ECD-rFc protein generated in 293T cells and a His-HER-2-ECD protein that was produced in bacteria and purified over a nickel column. The Igk leader sequence contained in the vectors directs the fusion proteins to be secreted. The vectors encoding the fusion proteins were transiently transfected into 293T cells, and the corresponding fusion proteins were secreted into the media. Production of the secreted fusion proteins was confirmed using a sandwich ELISA, and the concentrations were determined by comparison to an established CD147-rFc standard (data not shown).

To demonstrate that the generated MAPcL-rFc proteins were designed to be folded into a native conformation, an ELISA analysis was performed using commercially available anti-Her-2 antibodies generated against either native (monoclonal antibody 3F32 and Herceptin) or denatured (monoclonal antibody 3F27) HER-2 protein. Two antigens consisting of the ECD of HER-2 were analyzed: the conformation-carrying HER-2-ECD-rFc protein produced in 293T cells and a His-HER-2-ECD protein that was produced in bacteria and purified over a nickel column. The anti-native HER-2 antibody (3F32) recognized the HER-2-ECD-rFc produced in 293T (Fig. 1A), but was unable to detect the purified His-HER-2-ECD protein produced in bacteria (Fig. 1B). Also, Herceptin was unable to detect the denatured His-HER-2-ECD protein purified from bacteria (Fig. 1B). However, a strong response was observed for Herceptin when HER-2-ECD-rFc protein was used as the antigen for the ELISA analysis (Fig. 1A). Although the 3F27 antibody generated against denatured HER-2 did not detect the HER-2-ECD-rFc protein (Fig. 1A), this antibody had a strong response to bacterial HER-2-ECD (Fig. 1B).

To confirm the specific recognition of native versus denatured epitopes by the purchased antibodies, flow cytometry was performed on unfixed SKBR3 cells, a breast cancer cell line known to have HER-2 amplification (38). Because surface HER-2 would retain its native confirmation on the
unfixed SKBR3 cells, the anti-HER-2 3F27 antibody, specific for denatured HER-2, was unable to detect surface HER-2 on the cell membrane of SKBR3 cells by flow cytometry (Fig. 1C). When anti-HER-2 3F32 antibody and Herceptin, both of which recognize conformational HER-2, were used for flow cytometry analysis, a large shift in fluorescence was observed indicated that the antibodies recognized HER-2 present on the membrane of the SKBR3 cells (Fig. 1C).

A binding competition assay was performed to verify that the conformation-carrying antigen ELISA was recognizing the MAPcL antigen specifically. Wells were precoated with anti-rabbit IgG followed by HER-2-ECD-rFc. Purchased HER-2-Fc and CD30-Fc purified chimeric proteins (R&D Systems) were serially diluted and added to a constant amount of Herceptin before addition to the wells. The reactions were developed after incubation with the secondary antibody.

**Screening of patients for autoantibodies using the conformation-carrying antigen ELISA**

Twenty MAPcL–rFc fusion antigens designed to contain their native conformation were generated by cloning the sequences encoding the ECD or secreted proteins 5' of the rFc sequence (see Table 1 for identity of all 20 antigens). The expression plasmids were individually transfected into 293T cells, and the MAPcL–rFc fusion proteins were secreted into the media. The 20 fusion proteins were quantitated by sandwich ELISA analysis (data not shown). To detect autoantibodies in plasma collected from patients, a conformation-carrying antigen ELISA was developed using the generated MAPcL–rFc antigens. To immobilize the MAPcL–rFc fusion proteins, anti-rabbit IgG was used to precoat the wells of a 96-well plate. The media from the transfected 293T cells, which contains the generated MAPcL–rFc fusion proteins designed to have native conformations, was used to precoat the wells of a 96-well plate. The media from the transfected 293T cells, which contains the generated MAPcL–rFc fusion proteins designed to have native conformations, was used to precoat the wells of a 96-well plate. The reactions were developed after incubation with the secondary antibody.
Table 3. Absorbance measurements of autoantibodies and their association with breast cancer

<table>
<thead>
<tr>
<th>Autoantibody</th>
<th>Normal mammogram (n = 200)</th>
<th>Breast cancer (n = 200)</th>
<th>P valuea</th>
<th>ORb</th>
<th>95% CI</th>
<th>Increase in c-statisticc</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD320</td>
<td>0.15 (0.12)</td>
<td>0.16 (0.12)</td>
<td>0.62</td>
<td>1.10</td>
<td>0.90–1.35</td>
<td>0.000</td>
<td>0.96</td>
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<td>EPHA2</td>
<td>0.13 (0.06)</td>
<td>0.16 (0.10)</td>
<td>0.0006</td>
<td>1.64</td>
<td>1.21–2.24</td>
<td>0.034</td>
<td>0.037</td>
</tr>
<tr>
<td>GFRA1</td>
<td>0.18 (0.06)</td>
<td>0.20 (0.08)</td>
<td>0.0081</td>
<td>1.28</td>
<td>1.03–1.59</td>
<td>0.013</td>
<td>0.32</td>
</tr>
<tr>
<td>IGFBP2</td>
<td>0.21 (0.12)</td>
<td>0.25 (0.13)</td>
<td>0.0006</td>
<td>1.39</td>
<td>1.10–1.75</td>
<td>0.030</td>
<td>0.050</td>
</tr>
<tr>
<td>CST2</td>
<td>0.17 (0.09)</td>
<td>0.20 (0.10)</td>
<td>0.0013</td>
<td>1.39</td>
<td>1.12–1.73</td>
<td>0.026</td>
<td>0.13</td>
</tr>
<tr>
<td>GAL1</td>
<td>0.17 (0.06)</td>
<td>0.20 (0.07)</td>
<td>&lt;0.0001</td>
<td>1.75</td>
<td>1.37–2.23</td>
<td>0.051</td>
<td>0.021</td>
</tr>
<tr>
<td>HER-2</td>
<td>0.13 (0.04)</td>
<td>0.15 (0.06)</td>
<td>&lt;0.0001</td>
<td>1.65</td>
<td>1.28–2.13</td>
<td>0.039</td>
<td>0.054</td>
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<tr>
<td>LAMC2</td>
<td>0.15 (0.05)</td>
<td>0.17 (0.08)</td>
<td>0.0007</td>
<td>1.47</td>
<td>1.16–1.88</td>
<td>0.025</td>
<td>0.13</td>
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<td>ANGPTL4</td>
<td>0.18 (0.05)</td>
<td>0.20 (0.06)</td>
<td>0.0001</td>
<td>1.57</td>
<td>1.24–1.99</td>
<td>0.041</td>
<td>0.032</td>
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<tr>
<td>DKK1</td>
<td>0.18 (0.10)</td>
<td>0.24 (0.11)</td>
<td>&lt;0.0001</td>
<td>1.77</td>
<td>1.40–2.24</td>
<td>0.060</td>
<td>0.0093</td>
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<tr>
<td>MUC1</td>
<td>0.14 (0.06)</td>
<td>0.18 (0.08)</td>
<td>&lt;0.0001</td>
<td>1.83</td>
<td>1.41–2.37</td>
<td>0.055</td>
<td>0.012</td>
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<td>SRP2</td>
<td>0.14 (0.07)</td>
<td>0.17 (0.08)</td>
<td>0.0007</td>
<td>1.53</td>
<td>1.23–1.92</td>
<td>0.029</td>
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<td>LRP10</td>
<td>0.14 (0.05)</td>
<td>0.15 (0.07)</td>
<td>0.0098</td>
<td>1.35</td>
<td>1.09–1.68</td>
<td>0.011</td>
<td>0.47</td>
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<tr>
<td>LRRC15</td>
<td>0.11 (0.04)</td>
<td>0.12 (0.05)</td>
<td>0.005</td>
<td>1.09</td>
<td>0.89–1.34</td>
<td>0.001</td>
<td>0.82</td>
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<tr>
<td>SPINT2</td>
<td>0.15 (0.07)</td>
<td>0.18 (0.09)</td>
<td>0.0022</td>
<td>1.40</td>
<td>1.13–1.74</td>
<td>0.018</td>
<td>0.31</td>
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<tr>
<td>SPIN2</td>
<td>0.14 (0.07)</td>
<td>0.17 (0.08)</td>
<td>&lt;0.0001</td>
<td>1.65</td>
<td>1.31–2.07</td>
<td>0.042</td>
<td>0.052</td>
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<tr>
<td>CD147</td>
<td>0.10 (0.05)</td>
<td>0.12 (0.06)</td>
<td>0.0039</td>
<td>1.43</td>
<td>1.15–1.78</td>
<td>0.016</td>
<td>0.38</td>
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<tr>
<td>CDH9</td>
<td>0.10 (0.04)</td>
<td>0.12 (0.04)</td>
<td>0.0033</td>
<td>1.43</td>
<td>1.14–1.79</td>
<td>0.014</td>
<td>0.40</td>
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<tr>
<td>GRN</td>
<td>0.12 (0.06)</td>
<td>0.13 (0.07)</td>
<td>0.019</td>
<td>1.16</td>
<td>0.94–1.43</td>
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<td>0.65</td>
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<td>SUSD2</td>
<td>0.12 (0.04)</td>
<td>0.13 (0.05)</td>
<td>0.0085</td>
<td>1.36</td>
<td>1.10–1.70</td>
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<td>0.38</td>
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</table>

NOTE: Data shown as mean (SD) of √OD – Background.

aDifferences between groups were tested using t tests. Significant Bonferroni adjusted P value < 0.05/20 = 0.0025 are shown in bold.
bOR (95% CI) for breast cancer prevalence per 1 SD increase in autoantibody was determined using logistic regression models adjusted for age, race, BMI, and current smoking status.
cChange in area under the ROC curve (i.e., c-statistic) was determined when autoantibody was added to the adjusted logistic regression models.
study design and adjusting for BMI, race, and current smoking status. The group with the best model fit (i.e., minimum AIC) contained the autoantibody responses against the following 7 antigens: ANGPTL4, DKK1, GAL1, MUC1, GFRA1, GRN, and LRRC15 (Table 4). Of these 7, only autoantibody responses against ANGPTL4, DKK1, MUC1, and GAL1 individually showed a significant increase in the area under the ROC curve when added to the base model (Table 3). In the fully adjusted logistic regression model including the group of antigens, current smoking had the largest OR (95% CI) of prevalent breast cancer OR = 7.88 (2.68–23.2); and BMI was also a significant risk factor OR = 1.09 (1.04–1.13) per 1 kg/m² increase (Table 4). GAL1 had an OR of 6.73 (3.42–13.3), so a patient was almost 7 times as likely to have breast cancer per 1 SD increase in autoantibody response against GAL1. The autoantibody responses against GFRA1 (OR = 0.41), GRN (OR = 0.55), and LRRC15 (OR = 0.32) all had inverse associations with odds of prevalent breast cancer when adjusted for responses against the other antigens (Table 4). Taken together, the autoantibody response against the group of 7 antigens increased the area under the ROC curve from 0.64 to 0.82 (P < 0.0001) and had the following diagnostic measures: sensitivity (72.9%), specificity (76.0%), and positive likelihood ratio (95% CI) 3.04 (2.34–3.94; Fig. 2). The model was also calibrated across risk deciles (Hosmer–Lemeshow, P = 0.13).

Because breast cancer is a heterogeneous disease, it is possible that the autoantibody response against a combination of antigens may categorize a subtype of breast cancer differently than analyzing all breast cancer subtypes as a whole. The breast cancer samples were grouped into individual breast cancer subtypes: invasive, in situ, ER positive, tumor maximum dimension >1 cm, lymph node involvement, and HER-2 positive. The ability to discriminate cases from controls in each subtype was tested using autoantibody reactivity against the 7-antigen combination in addition to age, BMI, race, and current smoking status (Fig. 2). The 7-antigen combination model performed similarly in all subtypes of breast cancer; the c-statistic was 0.81 to 0.85. Of the breast cancer subtypes, in situ tumors had the greatest area under the ROC curve (0.8520, P < 0.0001) when analyzed for autoantibody responses against the 7-antigen combination. The model was not calibrated when considering only those cancers with lymph node involvement because of 4 unexpected breast cancers with very low model probabilities (Hosmer–Lemeshow P = 0.0036).

Discussion

Early detection of breast cancer allows a physician to treat the initial stage of the disease before metastasis, thereby allowing for a higher rate of remission or long-term survival for the patient. Detecting the presence of autoantibodies generated against tumor proteins in the blood of patients would be an ideal method for breast cancer detection. However, the tumor antigens need to be identified before specific autoantibody responses in patients can be ascertained. We generated a library that encodes membrane and secreted proteins that are highly expressed in breast cancer and may elicit an immune response.

We have shown that antigen conformation alters antibody-binding affinity in our assay, and the detection of autoantibodies is limited by epitope conformation (Fig. 1). We used a robust sample set to develop the conformation-carrying ELISA consisting of 200 plasma samples collected from newly diagnosed patients with breast cancer before surgery, chemotherapy, or radiation treatment. In addition, plasma was collected from 200 age-matched subjects defined by a confirmed normal mammogram in the preceding 6 months (Table 2). All 400 plasma samples were screened individually for autoantibody response against 20 TAAs designed to contain their native conformation using ELISA. Four of the 20 TAAs analyzed in our assay have previously been reported to generate an antibody response in patients with breast cancer: MUC1 (39, 40), HER-2 (41), IGBP2 (15), and GRN (42). Detection of autoantibodies against 12 of the 20 antigens was statistically significant for discriminating between normal and cancer samples (Table 3, bold). However, we did not observe a significant autoantibody response against GRN in our assay. Of the 12 significant antigens, 9 have not been previously associated with breast cancer autoantibodies. To our knowledge, this is the first report of the detection of autoantibodies against ANGPTL4, CST2, DKK1, EPHA2, GAL1, LAMC2, SPINT2, SPON2 and SSR2 in patients with breast cancer (Table 3).

Previously it has been shown that screening serum against a panel of antigens to detect autoantibodies compared with only a single antigen increases the sensitivity of the assay (17). This finding is consistent with the fact that breast cancer is a heterogeneous disease (43), and each individual patient’s immune system is distinct. A combination of 7 TAAs, consisting of ANGPTL4, DKK1, GAL1, MUC1, GFRA1, GRN, and LRRC15, had the greatest diagnostic capability (Table 4). Compared with previously published

Table 4. Multivariable logistic regression model odds ratios for breast cancer

<table>
<thead>
<tr>
<th>Variable</th>
<th>OR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (per 1 year)</td>
<td>1.00⁹ (0.98–1.02)</td>
</tr>
<tr>
<td>White race</td>
<td>0.70 (0.19–2.68)</td>
</tr>
<tr>
<td>BMI (per 1 kg/m²)</td>
<td>1.09 (1.04–1.13)</td>
</tr>
<tr>
<td>Current smoking</td>
<td>7.88 (2.68–23.2)</td>
</tr>
<tr>
<td>ANGPTL4 (per 1 SD)</td>
<td>1.71 (1.16–2.50)</td>
</tr>
<tr>
<td>DKK1 (per 1 SD)</td>
<td>1.87 (1.28–2.73)</td>
</tr>
<tr>
<td>GAL1 (per 1 SD)</td>
<td>6.73 (3.42–13.3)</td>
</tr>
<tr>
<td>GFRA1 (per 1 SD)</td>
<td>0.41 (0.21–0.82)</td>
</tr>
<tr>
<td>GRN (per 1 SD)</td>
<td>0.55 (0.38–0.81)</td>
</tr>
<tr>
<td>LRRC15 (per 1 SD)</td>
<td>0.32 (0.19–0.55)</td>
</tr>
<tr>
<td>MUC1 (per 1 SD)</td>
<td>1.67 (1.16–2.41)</td>
</tr>
</tbody>
</table>

⁹Because of individual 1:1 age matching.
Figure 2. ROC curve comparison for classification of patients with breast cancer. The autoantibody responses against 7 antigens (i.e., ANGPTL4, DKK1, GAL1, GFRA1, GRN, LRRC15, and MUC1) were added to a logistic regression model that included age, BMI, race, and current smoking status. The ROC curves were determined for all subjects (top) and by specific subtypes of breast cancer including ER positive, invasive, maximum tumor dimension >1 cm, in situ, lymph node involvement, and HER-2 amplification (bottom).
multiple antigen panels used to detect breast cancer autoantibodies (17, 44–46), the combination of these 7 TAs is unique, and our study contains the largest patient population of breast cancer and healthy samples. Interestingly, in the 7-antigen combination, 4 of the antigens have statistical significance individually (Table 3), but 3 of the antigens, GFRA1, GRN, and LRRC15, were not statistically significant on their own (Table 3). However, GFRA1, GRN, and LRRC15 were inversely associated with breast cancer, indicating that lower amounts of these autoantibodies in a patient, in combination with higher levels of the directly associated autoantibodies, increased the likelihood of having breast cancer (Table 4). When the 7 antigens were added to knowledge of current smoking status and BMI, the sensitivity and specificity of the assay was 72.9% and 76.0%, respectively. The area under the ROC curve (95% CI) was 0.82 (0.77–0.85), and the positive likelihood ratio was 3.04 for the conformation-carrying ELISA. Because breast cancer is a heterogeneous disease, patients were followed a patient for breast cancer recurrence (47). Comparing data, constructing databases): Writing, review, and/or revision of the manuscript: R.L. Evans, J.V. Pottala, K.A. Egland Study supervision: K.A. Egland

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References
Combination of Autoantibodies Discriminate Breast Cancer

Classifying Patients for Breast Cancer by Detection of Autoantibodies against a Panel of Conformation-Carrying Antigens

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