Computer-Generated Nuclear Features Compared With Axillary Lymph Node Status and Tumor Size as Indicators of Breast Cancer Survival

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The extent to which malignant cells deviate from normal is generally accepted to be a prognostic indicator. However, assessing the degree of deviation has been subjective and poorly reproducible. Our goal is to develop a computer program for objectively measuring nuclear size, shape, and texture from histologic slides and to make the program available on the Internet. We used this program to analyze 353 histologic sections obtained from patients with invasive breast cancer who were diagnosed and treated from 1981 through 1995 and who had determinable outcomes. The median follow-up was 8.3 years. We compared the relationship of survival with our computer-derived nuclear features versus axillary lymph node status and tumor size. We believe that our results are generally applicable because our patient survival, when stratified by lymph node status, was similar to that of the 24,000 breast cancer patients in the National Cancer Institute’s Surveillance, Epidemiology, and End Results program. In multivariate analysis, the strongest prognostic factor was the largest nuclear area, followed by tumor size and the extent of axillary lymph node involvement. The mean area of the 3 largest nuclei when combined with tumor size identified 30% of all breast cancer patients who had an 87% 15-year breast cancer-specific survival. Inclusion of lymph node status added little to this 2-factor model. Routine axillary lymph node surgery for prognostic purposes may become unnecessary, because nuclear features may provide sufficient information.

In this study we examined the relationship between breast cancer survival, the classical prognostic criteria of tumor size and lymph node status, and computer-generated nuclear features in histologic sections. We used a semiautomatic program that we developed and call Xcyt (available free of charge at http://dollar.biz.uiowa.edu/~street/xcyt/xcyt.html). No special tissue processing is required; thus others can verify our results with their own samples.

Previously, we used Xcyt to analyze fine-needle aspirates of breast cancers. In those cytologic samples, the nuclear feature of largest perimeter was prognostically stronger than either lymph node status or tumor size.

MATERIALS AND METHODS

Patients

Six hundred and twenty-nine patients with invasive breast cancer and without distant metastases underwent surgery from April 1981 through 1995. All surgery was done at the University of Wisconsin Hospital, and all cases were diagnosed by a University pathologist and reviewed according to quality assurance standards. Three hundred and fifty-three patients had determinable outcomes and tumor tissue available for resectioning. In each case, the histologic diagnosis was the one originally made. The diagnostic blocks were retrieved and recut. Most of the patients excluded from the study had undergone excisional biopsy at another facility but had axillary surgery with negative nodes at our institution. Material was not available for resectioning in such patients. For this reason, only 155 of the 333 node-negative (N−) patients could be included in this study, compared with 200 of the 262 node-positive (N+) patients. The mean tumor size was 2.7 cm for patients unavailable for follow-up, compared with 2.8 cm for those included in the study. For study patients, the median follow up was 8.3 years; 25 patients were followed for at least 15 years. Three hundred and thirty-seven patients underwent axillary lymph node dissections and pathologic staging. The 16 patients who did not undergo axillary surgery were clinically N− and had either small tumors of favorable histology or complicating medical conditions. None of the clinically staged patients subsequently developed axillary recurrences or distant metastases.

Our patient population is representative of the general breast cancer patient population of the United States. When stratified by axillary lymph node involvement, our patients’ survival was similar to that of 24,000 similar patients from the National Cancer Institute’s Surveillance, Epidemiology, and End Results (SEER) program. Survival was superimposed for the first 5 years and deviated only slightly thereafter (Fig 1). The differences between 5 and 10 years is not surprising because of the small number of our patients who had positive nodes and survived beyond 5 years and because of the decreasing prognostic importance of lymph node status with time.

Sample Preparation

Tissues were fixed in 10% neutral formalin, embedded in paraffin, and cut into 6-μm sections. These sections were stained with Weigart’s hematoxylin without counterstaining.
Selection of Nuclei for Analysis

A single, maximally atypical field was visually selected for analysis. A 640 × 480 digital image of the field was made using a ×63 objective on a microscope with a ×2.5 ocular on the video camera port. A frame grabber board (WinVision Pro; Quanta, Mountain View, CA) was used to capture the field as a .gif image. At this magnification, 3.9 pixels equal 1 micron in 1 dimension and 15.21 pixels equal 1 square micron in 2 dimensions. Analysis was performed on 10 to 100 nuclei per patient, depending on the number of nuclei present in the field.

Automatic Identification of Nuclei

The Xcyt program automatically locates and outlines cell nuclei in an image by searching for objects that match 1 of a collection of elliptical templates. The templates are updated as the system is used to reflect the shape of the objects found in previous images. In this way, the search becomes both faster and more accurate as the system is used. (Street\(^8\) has provided more details on the image analysis performed by Xcyt.) On average, approximately 75% of the nuclei in the images are correctly outlined. Erroneous outlines are edited or deleted by the user, and missed nuclei are traced manually.

Nuclear Morphometric Features

The following nuclear features were computed for each identified nucleus:\(^2\):
- **Radius** was computed by averaging the length of radial line segments from the center of the nuclear mass to each of the points of the nuclear border.
- **Perimeter** was measured as the distance around the nuclear border.
- **Area** was calculated by counting the number of pixels in the interior of the nuclear border, then adding half of the pixels on the perimeter.
- Perimeter and area were combined to give a measure of the compactness of the cell nuclei using the formula \(\frac{\text{perimeter}^2}{\text{area}}\).
- **Smoothness** was quantified by measuring the difference between the length of each radius and the mean length of adjacent radii.
- **Concavity** measured the size of any indentations in the nuclear border.
- **Concave points** counted the number of points on the nuclear border that lie on an indentation.
- **Symmetry** was measured by finding the relative difference in length between line segments perpendicular to and on either side of the major axis.
- **Fractal dimension** was approximated using the “coastline approximation” described by Mandelbrot,\(^9\) which measures nuclear border irregularity.
- **Texture** was measured by finding the variance of the gray scale intensities in the component pixels.

For these 10 features, the computer calculated the mean value, the “largest” value, and the standard error for each nuclear feature, resulting in a total of 30 features. The largest value for each feature was the mean of the 3 largest values for all nuclei in the analyzed image; 3 was chosen as the smallest number that would guard against numerical instability in the shape features.

Statistical Analysis

Calculations were done using SPSS software (SPSS, Chicago, IL).\(^10,11\) The first step was to use the Cox univariate analysis to rank the available predictive factors according to their individual ability to predict time of breast cancer death. The independent variables were the 30 nuclear features, along with tumor size and lymph node status. Patients were then assigned to groups based on splits for each pair of the 3 strongest prognostic factors: tumor size, largest nuclear area, and lymph node positivity. Splits were at the median for tumor size and largest area and positive or negative for axillary lymph node status.
TABLE 1. Comparison of Prognosis for Patients Divided Into Groups Based on Largest Nuclear Area of Their Cancers

<table>
<thead>
<tr>
<th>Nuclear Largest Area Cutpoint ($\mu^2$)</th>
<th>$P$</th>
<th>Relative Risk Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>45.6</td>
<td>.0200</td>
<td>3.9</td>
</tr>
<tr>
<td>58.4</td>
<td>.0009</td>
<td>4.1</td>
</tr>
<tr>
<td>67.6</td>
<td>.0007</td>
<td>2.7</td>
</tr>
<tr>
<td>78.0</td>
<td>.0004</td>
<td>2.4</td>
</tr>
<tr>
<td>87.1</td>
<td>.0757</td>
<td>1.8</td>
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<tr>
<td>100.5</td>
<td>.0004</td>
<td>2.1</td>
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<tr>
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<td>.0002</td>
<td>2.2</td>
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<tr>
<td>137.0</td>
<td>.0004</td>
<td>2.3</td>
</tr>
<tr>
<td>172.3</td>
<td>.0094</td>
<td>2.1</td>
</tr>
</tbody>
</table>

NOTE. Cox regression analysis of breast cancer survival was performed for patients with cancers with larger nuclear largest areas than the cutpoints listed here compared with patients with having smaller ones.

RESULTS

The following nuclear features were highly related to breast cancer-specific overall survival in univariate analysis ($P < .0001$): mean area, mean perimeter, mean radius, standard error of area, largest area, and largest perimeter. Nuclear features of lesser significance ($P$ between .0001 and .01) were mean fractal dimension, standard error of perimeter, standard error of radius, and largest texture. Additionally, the number of cancerous axillary lymph nodes, and tumor size were highly significant with $P < .0001$. Overall, the 3 most significant factors were tumor size, axillary lymph nodes, and largest nuclear area. In a 3-factor Cox multivariate analysis model, the significance of largest area was $P < .0001$, tumor size was $P = .0001$, and N+ or N− was $P = .0171$. To determine the robustness of largest area as a prognostic factor, the patients were divided into deciles according to largest area. At all deciles, patients whose tumors had larger largest area nuclei had a poorer prognosis than those with smaller ones (Table 1). In comparison, N+ versus N− was $P = .0002$ with a relative risk of 2.5, and tumor size ≥2.4 cm versus tumor size <2.4 cm was $P < .0001$ with a relative risk of 3.8. We used the median cutpoint for largest area, even though the poorest prognostic separation occurred at the median cutpoint. Other cutpoints may produce better results.

Patients were divided according to the median values for largest area [87.1 $\mu^2$ (1325 pixels)], for tumor size (2.4 cm), and according to N+ or N− status. At these cutpoints, Kaplan-Meier survival analysis showed that patients were best separated according to whether their tumor size was smaller or larger than the median size, next best separated according to whether they were N+ or N−, and least well separated by whether their tumor’s nuclear largest area was smaller or larger than the median large area. The log rank statistics were 32.8 ($P < .0001$), 15.2 ($P = .0001$), and 7.9 ($P = .0049$), respectively. These 3 features were then paired according to the aforementioned splits to create 4 groups of patients for each pair. Taken in pairs, the overall $\chi^2$ value in Cox multivariate models was 50.0 for tumor size and largest area, 47.2 for largest area and lymph nodes, and 42.3 for tumor size and lymph nodes. Because the pair of tumor size and largest area gave the best separation, further studies were done with this pair; specifically, size <2.4 cm and largest area <87.1 $\mu^2$ (107 patients), size ≥2.4 cm and largest area ≥87.1 $\mu^2$ (67 patients), size ≥2.4 cm and largest area <87.1 $\mu^2$ (67 patients), and size ≥2.4 cm and largest area ≥87.1 $\mu^2$ (106 patients). Within these 4 groups, Kaplan-Meier analysis revealed no significant difference between breast cancer survival of N+ and N− patients, and no significant differences were noted in the number of patients with distant disease recurrence and in breast cancer survival (Student’s $t$ test). The recurrences of distant disease in these 4 groups are displayed in Figure 2.

The 15-year breast cancer-specific survival for these 4 groups was 86.6%, 77.9%, 35.4%, and 48.9%, respectively. All group separations were significant except when the size of the cancer was ≥2.4 cm, in which case the nuclear largest area was insignificant (Table 2). Kaplan-Meier survival curves are shown in Figure 3, and the significance of the difference between the curves is delineated in Table 2.

DISCUSSION

Several researchers, beginning with Black et al., showed that cellular features were related to outcome. Subsequently, various grading systems were proposed. However, nuclear grade is rarely used as a prognostic indicator because of the subjectivity inherent in its assessment. Instead, decisions regarding postopera-

![Figure 2](image-url)
tive treatment regimens are typically based primarily on the tumor size, lymph node, metastasis (TNM) staging system. Determination of axillary node status for TNM staging necessitates surgical removal of axillary lymph nodes. Complete removal leaves the patient with numbness and other complications, including a 15% risk of arm lymphedema. Less-extensive dissections, such as sentinel lymph node sampling, are promising but not yet clinically established approaches. Thus the focus of our research has been on investigating the prognostic significance of objectively defined nuclear features and to determine whether the breast cancer staging without using lymph node information is feasible.

Other computer-based studies have demonstrated that larger nuclear size portends a poorer prognosis. Our largest area values were about 70% greater than the mean area values (101 ± 56 μ² versus 60 ± 28 μ²) and probably identified multiploid nuclei that have been shown by both flow cytometry and DNA labeling index to be associated with an unfavorable prognosis.

In our patients, tumor size was a stronger prognostic feature than axillary lymph node status, yet other studies found lymph node status to be the stronger feature. Our patient population differed from older series, because our study included small cancers that were identified either by either aggressive fine-needle aspiration or mammography. For instance, 2.54% of our patients had tumors smaller than 0.5 cm compared with 0.78% of patients in the SEER database. We found that N⁺ versus N⁻ was the second to tumor size in strength as a prognostic factor, yet largest area was stronger in multivariate analysis. This is explained because tumor size and N⁺ or N⁻ are closer correlated than tumor size and large area (Pearson correlation coefficient of 0.459 versus 0.199). Therefore, largest area adds more information to tumor size than does axillary lymph node status. Within the 4 groups based on tumor size and largest area, we were unable to demonstrate significant breast cancer survival differences between N⁺ and N⁻ patients. Yet nodal positivity was statistically significant (P = 0.0171) in the 3-factor Cox model. Further studies with more patients are needed to determine whether additional clinically useful prognostic information can be obtained from axillary lymph node examination.

In previous cytologic analyses of breast fine-needle aspirates, we found that the nuclear size feature of largest perimeter was the strongest prognostic nuclear feature for overall survival. This is consistent with our current results, because largest perimeter and largest area are strongly correlated (Pearson correlation coefficient, 0.957, P < .001). The only inconsistency between our cytologic and histologic findings lies in the relative importance of nuclear size and tumor size above the median cutoffs. Cytologically, large and small-sized cancers had a similarly poor prognosis when associated with unfavorable cytology. Histologically, favorable or unfavorable cytology was important only in small-sized cancers and made no difference in larger ones. To try to understand this inconsistency, we compared cytology and histology in the 136 patients for whom both types of material was available. Surprisingly, the mean values for largest perimeter and for largest area were virtually identical in both types of preparations. The correlation...
(Pearson) between cytologic largest perimeter and histologic largest area was 0.54 \((P < .001)\). Although highly statistically significant, there were sufficient differences in individual patients to explain the inconsistency between cytologic and histologic results.

Our study’s endpoint was breast cancer–specific survival rather than either distant disease-free survival or overall survival for 3 reasons. First, our study was not controlled for the use of adjuvant chemotherapy. Axillary lymph node–positive patients were generally given adjunctive chemotherapy, whereas N– patients were not. The protocol adjunctive chemotherapy, mainly cyclophosphamide, methotrexate, 5-fluorouracil (CMF)–based, that was given to our N+ patients was shown to prolong distant disease-free survival but to not increase breast cancer–specific survival.\(^{31-35}\) Second, death from causes unrelated to breast cancer was common because of the long follow-up and the preponderance of elderly patients. Third, the time of recurrence was difficult to document during long-term follow-up because patients were followed at inconsistent intervals after 5 years. Nonetheless, similar results were obtained when distant disease- free survival rather than overall breast cancer–specific survival was used as the endpoint.

Only modest time and money investments are required to add Xcyt to a pathologist’s practice. Sample preparation is similar to that used for routine histologic examination, except that Weigart’s hematoxylin is used to accentuate nuclear membranes and no counterstain is applied. Images can be captured with either a video or digital still camera mounted on a microscope. In many instances, an existing 35-mm film camera can be replaced by a digital single-lens reflex camera, and the acquired image files can be transferred to a computer without a video camera or a frame-grabber board. A maximally atypical field can be selected for analysis during the course of routine histologic examination. The Xcyt analysis takes about 15 minutes. Xcyt automatically outlines the first 5 nuclei relatively slowly but speeds up markedly for subsequent nuclei. After automatic outlining, an operator edits errant outlines and manually outlines missed nuclei. Nuclear feature calculations are virtually instantaneous.

We conclude that the prognostic importance of tumor size and axillary lymph node status needs to be evaluated in conjunction with computer-based image analysis. Currently, the major reason for axillary lymph node staging is to determine the need for adjunctive chemotherapy and the type of this therapy indicated. Tumor size together with the information obtained via image analysis may provide comparable prognostic information and thus obviate the need for routine axillary lymph node surgery.

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REFERENCES
11. SPSS Advanced Statistics 7.5. Chicago, IL, SPSS Inc., 1997


