Introduction: Tumour location within the breast varies with the highest frequency in the upper outer quadrant (UOQ) and lowest frequency in the lower inner quadrant (LIQ). Whether tumour location is prognostic is unclear. To determine whether tumour location is prognostic, associations between tumour site and clinicopathological characteristics were evaluated.

Materials and Methods: All patients enrolled in the Clinical Breast Care Project whose tumour site—UOQ, upper inner quadrant (UIQ), central, LIQ, lower outer quadrant (LOQ)—was determined by a single, dedicated breast pathologist were included in this study. Patients with multicentric disease (n = 122) or tumours spanning multiple quadrants (n = 381) were excluded from further analysis. Clinicopathological characteristics were analysed using chi-square tests for univariate analysis with multivariate analysis performed using principal components analysis (PCA) and multiple logistic regression. Significance was defined as $P < 0.05$.

Results: Of the 980 patients with defined tumour location, 30 had bilateral disease. Tumour location in the UOQ (51.5%) was significantly higher than in the UIQ (15.6%), LOQ (14.2%), central (10.6%), or LIQ (8.1%). Tumours in the central quadrant were significantly more likely to have higher tumour stage ($P = 0.003$) and size ($P < 0.001$), metastatic lymph nodes ($P < 0.001$), and mortality ($P = 0.011$). After multivariate analysis, only tumour size and lymph node status remained significantly associated with survival.

Conclusions: Evaluation of tumour location as a prognostic factor revealed that although tumours in the central region are associated with less favourable outcome, these associations are not independent of location but rather driven by larger tumour size. Tumours in the central region are more difficult to detect mammographically, resulting in larger tumour size at diagnosis and thus less favourable prognosis. Together, these data demonstrate that tumour location is not an independent prognostic factor.

Keywords: breast quadrants, tumour location, prognosis
Background

Although mortality rates from breast cancer have decreased since 1990, it is estimated that >39,500 women in the United States died from breast cancer in 2013 [1]. A number of pathological characteristics have been identified, which can be used as prognostic factors, including tumour size and grade, hormone receptor, HER2, and lymph node status [2, 3], which are all considered when predicting prognosis and determining the most effective treatment options. In addition, molecular tests have been developed that add prognostic value by determining intrinsic subtype, stratification of tumours into low or high grade, or predicting recurrence [4–7]. Despite these factors and tests, predicting patient outcome is imprecise, as patients with similar pathological characteristics treated with identical regimens have highly variable clinical outcomes [8], suggesting that identification of additional prognostic factors may improve patient stratification.

Tumour location within the breast has been proposed as an independent prognostic factor. For example, the frequency of axillary lymph node metastasis was significantly lower in the upper inner quadrant (UIQ, 20.6%) compared to all other quadrants (33.2%) [9]. In contrast, tumours in the upper outer quadrant (UOQ), which is the most frequent site of tumour location, have been associated with improved survival compared to other quadrants [10, 11]; survival data for non-UOQ tumours have been mixed with findings demonstrating decreased survival for the lower inner quadrant (LIQ), and lower, medial or periareolar regions [12–17]. In contrast, other studies have found no association between tumour location and outcome [18–20].

Reasons for discordant results are varied. For example, the manner in which tumour locations are grouped can affect survival results: when quadrants were evaluated individually, patients with tumours in the LIQ had a twofold increase in mortality; however, using the same data set, when tumour sites were combined into inner versus outer or lower versus upper, survival did not differ [13]. Other studies have suggested that the less favourable outcomes associated with tumours in the medial region can be attributed to undetected positive lymph nodes in internal mammary lymph nodes [12, 13], although several studies have demonstrated that treatment of the internal mammary lymph nodes does not improve survival [20–23]. Finally, tumour locations demonstrating less favourable survival are associated with known prognostic factors such as increased tumour size, grade, or lymph node status [13, 16]. Thus, to determine whether tumour location is an independent prognostic factor, data from the Clinical Breast Care Project (CBCP) was analysed to determine whether tumour site was associated with any clinicopathological characteristics.

Methods

Patient enrolment and consent

For inclusion in the CBCP, all patients met the following criteria: (1) adult over the age of 18 years, (2) mentally competent and willing to provide informed consent, and (3) presenting to the breast centres with evidence of possible breast disease. Tissues were collected with approval from the Walter Reed National Military Medical Centre (WRNMMC) Human Use Committee and Institutional Review Board. All enrolled subjects voluntarily agreed to participate and were provided with layered consent forms that included permission to gather tissue and that described the primary research uses of the samples.

Pathological characterisation

The CBCP database was queried to identify all patients with known tumour location diagnosed 2001–2013. Tumour location was classified using the SEER Coding guidelines (http://seer.cancer.gov/archive/manuals/2010/AppendixC/breast/coding_guidelines.pdf). Pathological characterisation of all specimens was performed by a single, dedicated breast pathologist as previously described [24]. To ensure consistency, diagnosis of all tumour samples were made by one pathologist from haematoxylin and eosin stained slides; staging was performed using guidelines defined by the AJCC Cancer Staging Manual seventh edition [25] and grade was assigned using the Bloom–Richardson system of classification [26, 27]. Oestrogen receptor (ER), progesterone receptor (PR), and HER2 status
were determined by immunohistochemical (IHC) analysis (MDR Global, Windber, PA); the cut-off for defining ER and PR positivity was ≥1% positively staining cells following the ASCO/CAP guidelines [28] while cases with HER2 scores = 2+ were then evaluated by fluorescence in situ hybridisation using the PathVysion® HER-2 DNA Probe kit (Abbott Laboratories, Abbott Park, IL).

Statistical Analysis

Univariate analysis was performed using chi-square analysis (http://www.physics.csbsju.edu/stats/contingency_NROW_NCOLUMN_form.html). After initial univariate analysis revealed highly associated variables, a multivariate approach was used to determine the extent to which each could be considered an independent prognostic factor. In the initial multivariate analysis, seven possible predictor variables were included: lymph node status, PR status, ER status, HER2 status, location, tumour grade, and size. Principal components analysis (PCA) was performed in order to reduce the dimensionality of a subsequent regression model and to create uncorrelated variables. Multiple logistic regression analysis was performed to assess the influence of these components on patient survival. The informative components were then decomposed to determine the true independent predictors of patient survival resulting from the regression. Further multiple logistic regression analysis was performed on lymph node status, tumour size, and tumour grade in order to determine how strongly other characteristics were associated with these variables.

Because our variables were categorical, one category for each was established as the reference point to which the other categories were compared in regression modelling. The reference categories corresponded to a negative status within the lymph node, PR, ER, and HER2 variables, the central region for tumour location, tumour grade 1, and tumour size T1. The odds ratios established through regression for the complementary categories represent the relative odds that a change from the reference category to the alternative category would cause a change in the dependent variable when all other variables are held constant. Statistical significance was defined as \( P < 0.05 \). Statistical analysis was carried out using R version 3.1.1 (http://www.R-project.org/).

Results

Patient characteristics

As of July 2014, 1,483 patients were enrolled in the CBCP. Of these, 122 were diagnosed with multicentric disease, and 381 had tumour locations spanning multiple quadrants (e.g., mid-inner), leaving 980 eligible patients for this study. The average age at diagnosis was 58.7. The majority of patients were Caucasian (80%), and the majority of tumours were early stage (88%), ER+HER2− (72%), T1 (66%) with negative lymph nodes (70%). Tumour grade was split into well-differentiated (29%), moderately differentiated (38%) and poorly differentiated (33%). Twenty-nine patients have died of disease (Figure 1). Median follow-up by tumour location was 5.3 years (range 1.6–13.8) for UOQ, 5.4 years (range 1.2–13.2) for UIQ, 5.6 years (range 2.0–13.2) for LOQ, 5.8 years (range 1.9–13.3) for LIQ, and 5.6 years (range 1.7–13.9) for central.

Differences in pathological characteristics by tumour location

Tumour location was higher in the UOQ (51.5%) compared to the UIQ (15.6%), lower outer quadrant (LOQ, 14.2%), central (10.6%), or LIQ (8.1%). To evaluate whether breast cancer risk factors are associated with tumour location, age at diagnosis, ethnicity, body mass index (BMI), and breastfeeding were evaluated by quadrant (Table 1); none of these factors differed significantly. Analysis of pathological factors revealed significant differences for tumour stage, size, lymph node status, and survival (Table 2), with central quadrants having a higher frequency of late-stage tumours (25%) compared to the other quadrants (range 7–13%), T3 tumours (16%) compared to other quadrants (range 0–5%) and metastatic lymph nodes (51%) compared to other quadrants (range 24–38%). Breast cancer mortality rates were highest in patients with tumours in the central quadrant (7%) compared to other quadrants (range 1–5%).
Figure 1. Survival curve by tumour location. Median time to death was 1.52 years for UOQ, 5.69 years for UIQ, 3.08 years for LOQ, 6.03 years for LIQ, and 1.77 years for central.

Table 1. Comparison of risk factors at diagnosis between tumour locations. Data are presented as proportion of individuals within each category.

|                      | UOQ (n = 520) | UIQ (n = 158) | LOQ (n = 143) | LIQ (n = 82) | Central (n = 107) | P value |
|----------------------|---------------|---------------|---------------|--------------|-------------------|---------|
| Age at Diagnosis     |               |               |               |              |                   |         |
| <40 years            | 0.06          | 0.04          | 0.05          | 0.07         | 0.03              | 0.115   |
| 40–49 years          | 0.20          | 0.22          | 0.29          | 0.12         | 0.20              |         |
| ≥50 years            | 0.74          | 0.74          | 0.66          | 0.81         | 0.77              |         |
| Menopausal status    |               |               |               |              |                   |         |
| Pre-menopausal       | 0.29          | 0.28          | 0.36          | 0.25         | 0.23              | 0.190   |
| Post-menopausal      | 0.71          | 0.72          | 0.64          | 0.75         | 0.77              |         |
| Ethnicity            |               |               |               |              |                   |         |
| African American     | 0.20          | 0.21          | 0.17          | 0.21         | 0.13              | 0.420   |
| Asian                | 0.01          | 0.03          | 0.01          | 0.01         | 0.03              |         |
| Hispanic             | 0.01          | 0.01          | 0.01          | 0.01         | 0.04              |         |
| Other                | 0.01          | 0.01          | 0.01          | 0.00         | 0.00              |         |
| Caucasian            | 0.77          | 0.74          | 0.80          | 0.77         | 0.80              |         |
| BMI                  |               |               |               |              |                   |         |
| <18.5                | 0.01          | 0.01          | 0.01          | 0.01         | 0.01              | 0.247   |
| 18.5–24.9            | 0.32          | 0.24          | 0.31          | 0.25         | 0.38              |         |
| 25–29.9              | 0.29          | 0.36          | 0.27          | 0.34         | 0.37              |         |
| ≥30                  | 0.38          | 0.39          | 0.41          | 0.40         | 0.24              |         |
| Breastfeed\*         |               |               |               |              |                   |         |
| Yes                  | 0.46          | 0.44          | 0.47          | 0.53         | 0.49              | 0.616   |
| No                   | 0.54          | 0.56          | 0.53          | 0.47         | 0.51              |         |

*Breastfeeding was considered only in parous women.
Table 2. Pathological characteristics at diagnosis by tumour location. Data are presented as proportion of individuals within each category.

|                      | UOQ (n = 520) | UIQ (n = 158) | LOQ (n = 143) | LIQ (n = 82) | Central (n = 107) | \(P\) value |
|----------------------|---------------|---------------|---------------|--------------|------------------|-------------|
| **Stage**            |               |               |               |              |                  |             |
| I                    | 0.56          | 0.66          | 0.53          | 0.59         | 0.46             | 0.003       |
| II                   | 0.33          | 0.27          | 0.34          | 0.30         | 0.29             |             |
| III                  | 0.09          | 0.06          | 0.11          | 0.06         | 0.18             |             |
| IV                   | 0.02          | 0.01          | 0.02          | 0.05         | 0.07             |             |
| **ER/HER2 status**   |               |               |               |              |                  | 0.124       |
| ER+/HER2−             | 0.70          | 0.76          | 0.68          | 0.76         | 0.71             |             |
| ER+/HER2−             | 0.15          | 0.08          | 0.10          | 0.05         | 0.07             |             |
| ER−/HER2+             | 0.05          | 0.10          | 0.07          | 0.04         | 0.06             |             |
| ER−/HER2−             | 0.10          | 0.06          | 0.15          | 0.15         | 0.16             |             |
| **PR Status**         |               |               |               |              |                  | 0.248       |
| Positive              | 0.62          | 0.71          | 0.67          | 0.63         | 0.64             |             |
| Negative              | 0.38          | 0.29          | 0.33          | 0.37         | 0.36             |             |
| **Lymph node status** |               |               |               |              |                  | <0.001      |
| Negative              | 0.64          | 0.76          | 0.62          | 0.66         | 0.49             |             |
| Positive              | 0.36          | 0.24          | 0.38          | 0.34         | 0.51             |             |
| **Tumour Grade**      |               |               |               |              |                  | 0.668       |
| Low                   | 0.29          | 0.33          | 0.27          | 0.26         | 0.27             |             |
| Intermediate          | 0.37          | 0.33          | 0.41          | 0.43         | 0.45             |             |
| High                  | 0.34          | 0.34          | 0.32          | 0.31         | 0.28             |             |
| **Ki67**              |               |               |               |              |                  | 0.174       |
| <20%                  | 0.58          | 0.55          | 0.50          | 0.42         | 0.50             |             |
| ≥20%                  | 0.42          | 0.45          | 0.50          | 0.58         | 0.50             |             |
| **Tumour Size**       |               |               |               |              |                  | <0.001      |
| T1                    | 0.67          | 0.71          | 0.70          | 0.71         | 0.52             |             |
| T2                    | 0.28          | 0.25          | 0.27          | 0.29         | 0.32             |             |
| T3                    | 0.05          | 0.04          | 0.03          | 0.00         | 0.16             |             |
| **Angiolymphatic invasion** |           |               |               |              |                  | 0.272       |
| Absent                | 0.77          | 0.78          | 0.73          | 0.79         | 0.68             |             |
| Present               | 0.23          | 0.22          | 0.27          | 0.21         | 0.32             |             |
| **Dead of Disease**   |               |               |               |              |                  | 0.011       |
| Yes                   | 0.02          | 0.01          | 0.05          | 0.05         | 0.07             |             |
| No                    | 0.98          | 0.99          | 0.95          | 0.95         | 0.93             |             |

Given the linear relationship between tumour size and lymph node involvement [29], lymph node status was compared within tumour size groups T1, T2, or T3. The frequency of positive lymph nodes did not differ significantly by tumour location for any of the sizes. Multivariate analysis was performed using all of the pathological characteristics; only lymph node status and tumour size remained significant predictors of breast cancer mortality with lymph node metastases and T3 tumour size having expected odds ratios of 3.5 (\(P = 0.026\)) and
5.1 \( (P = 0.019) \), respectively. Tumour size was significantly associated with positive lymph node status with T2 and T3 tumours having expected odds ratios of 12.3 \( (P < 0.001) \) and 24.7 \( (P < 0.001) \); high-grade tumours were also associated with positive lymph node status with an expected odds ratio of 1.6 \( (P = 0.045) \). Tumour location was only associated with tumour size, with all quadrants having smaller tumour size compared to the central region (expected odds ratios range \( 0.30–0.41)\).

**Discussion**

Determination of whether tumour location can be used prognostically is important in optimising treatment. Tumour location is highest in the UOQ (50–58%) across multiple populations, including Chinese, Danish, the United Kingdom and women treated within the United States Department of Defence healthcare system \[10, 11, 14, 30\]. Two studies suggest that the frequency of tumours in the UOQ has increased over time \[30, 31\]; given the association of tumour in the UOQ with improved prognosis, these data would suggest a trend towards a reduction in breast cancer mortality. Tumour occurrence in the LIQ, however, has also increased significantly and tumour location in the LIQ has been associated with >2-fold increase in mortality \[13\], suggesting that the decrease in mortality associated with increasing tumour location in the UOQ may be offset by concomitant increases in the LIQ. In our study, tumours in the UOQ showed a trend towards favourable prognosis although this did not reach the level of significance \( (P = 0.0754) \).

The less favourable prognosis seen in patients with tumours in the central region can be attributed to increased tumour size and positive lymph node status. Tumour size is thought to reflect the chronological age of the tumour, with smaller tumours being resected earlier than larger tumours \[32\]. Tumour size has been associated with positive lymph node status in multiple studies \[29, 33–41\]. In addition, tumour size is also prognostic in patients with both negative and positive lymph node status \[42–46\]. Thus, tumour location in the central region is a surrogate for larger tumour size, resulting in increased rates of metastatic lymph node and breast cancer mortality.

Previous studies have demonstrated that tumours within the central region are harder to detect than at other sites \[47, 48\] and that tumours in this region are more easily detected by clinical examination than mammography \[49\]. This difficulty in detecting tumours in the central region have been attributed to overpenetration of X-rays in the nipple-areolar complex; accurate diagnosis in this region may require the use of multiple imaging modalities \[50\]. The difficulty in mammographic detection and more frequent occurrence of palpable tumours in the central region explains the larger tumour size, and hence increased mortality found in our study.

Although this study included data from 980 patients, two other studies evaluated much larger data sets including tumour registry data from the US Department of Defence \( (n = 13,984) \) or Danish Breast Cancer Cooperative Group \( (n = 35,319) \). In both studies, UOQ was associated with more favourable prognosis, with one study reporting a hazard ratio of 0.820 and the other a 20% improved survival in patients whose tumours were in the UOQ \[10, 11\]. Our data demonstrated a trend towards improved prognosis for patients with tumours in the UOQ (expected odds ratio = 0.2988, \( P = 0.0754) \) when compared to the central region; this survival advantage may have reached the level of significance with a larger sample size. In addition, the median follow-up for these patients was 5.5 years; thus, differences in late recurrence \( (>5 \text{ years after diagnosis}) \) by tumour location may not have been detected. Finally, treatment data for each patient were not available. While some studies suggest that less favourable prognosis by tumour location has been associated with under-staging and suboptimal treatment of tumours in the LIQ when lymph node status of the internal mammary chain is not evaluated \[12, 15\], others studies found that treatment differences were not responsible for the decreased survival for patients with tumours in the LIQ \[13, 21, 22\]. Without these data in our study, it is not possible to determine whether the less favourable prognosis for patients with tumours in the central region, or conversely, the improved prognosis for those with tumours in the UOQ, reflects differential provision or response to treatment.

This study does have several advantages. Tumour location was specified by a single, dedicated breast pathologist, which provides a level of consistency not available in tumour registries or databases with pathological data compiled from multiple pathologists. In addition, our data were analysed by quadrant site, rather than grouped into lateral/medial \[14, 15\], which studies have shown may dilute the ability to detect associations between pathology and tumour location \[13\]. Finally, while we included tumours from the central region in this study, other studies did not include this region in statistical analysis \[11, 15\].
Conclusions

In this study, the UOQ, which was the most common site for tumours within the human breast, was not significantly associated with improved survival. In addition, although central tumour location was associated with higher mortality rates on univariate analysis, multivariate analysis demonstrated that centrality was not an independent risk for survival but rather, the association with less favourable prognosis was attributable to a higher frequency of T3 tumours. Difficulty in imaging and detecting tumours within the central region leads to delayed detection and larger tumour size at diagnosis, and tumour size ≥5 cm (T3) is an independent prognostic factor for both lymph node metastases and poor outcome. Together, these data suggest that tumour location is not prognostic.

Conflicts of interest

The author(s) declare that they have no conflict of interest.

Authors’ contributions

Seth Rummel collected the data and performed preliminary analyses; Nick Constantino performed multivariate analysis, Matthew T Hueman and Craig D Shriver provided clinical input and reviewed the manuscript, Rachel E Ellsworth conceptualised the study and drafted the manuscript.

Acknowledgments

The opinion and assertions contained herein are the private views of the authors and are not to be construed as official or as representing the views of the Department of the Army or the Department of Defence. This research was supported by a grant from the Office of the Congressionally Directed Medical Research Programs (Department of Defence Breast Cancer Research Program W81XWH-11-2-0135).

References

1. American Cancer Society: Breast Cancer Facts & Figures 2013–2014 In Atlanta: American Cancer Society, Inc 2013
2. Ellsworth RE, Decewicz DJ, Shriver CD and Ellsworth DL (2010) Breast cancer in the personal genomics era Curr Genomics 11(3) 146–161 DOI: 10.2174/138920210791110951 PMID: 21037853 PMCID: 2878980
3. Van de Vijer MJ (2014) Molecular tests as prognostic factors in breast cancer Virchows Arch 464 283–291 DOI: 10.1007/s00428-014-1539-0
4. Dowsett M, Sestak I, Lopez-Knowles E and Sidhu K et al (2013) Comparison of PAM50 risk of recurrence score with oncotype DX and IHC4 for predicting risk of distant recurrence after endocrine therapy J Clin Oncol 31(22) 2783–2790 DOI: 10.1200/JCO.2012.46.1558 PMID: 23816962
5. Paik S, Shak S and Tang G et al (2004) A multigene assay to predict recurrence of tamoxifen-treated, node-negative breast cancer N Engl J Med 351(27) 2817–2826 DOI: 10.1056/NEJMo041588 PMID: 15591335
6. Sotiriou C, Wirapati P and Loi S et al (2006) Gene expression profiling in breast cancer: understanding the molecular basis of histologic grade to improve prognosis J Natl Cancer Inst 98(4) 262–272 DOI: 10.1093/jnci/dji052 PMID: 16478745
7. Van de Vijer MJ, He YD and Van’t Veer LJ et al (2002) A gene-expression signature as a predictor of survival in breast cancer N Engl J Med 347(25) 1999–2009 DOI: 10.1056/NEJMoa021967

8. Gort M, Broekhuis M, Otter R and Klazinga NS (2007) Improvement of best practice in early breast cancer: actionable surgeon and hospital factors Breast Cancer Res Treat 102(2) 219–226 DOI: 10.1007/s10549-006-9327-4

9. Bevilacqua J, Cody H 3rd, MacDonald KA and Tan LK et al (2002) A prospective validated model for predicting axillary node metastases based on 2,000 sentinel node procedures: the role of tumour location [corrected] Eur J Surg Oncol 28(5) 490–500 DOI: 10.1053/ejso.2002.1268 PMID: 12217300

10. Kroman N, Wohlfahrt J, Mouridsen HT and Melbye M (2003) Influence of tumor location on breast cancer prognosis Int J Cancer 105(4) 542–545 DOI: 10.1002/ijc.11116 PMID: 12712447

11. Sohn VY, Arthurs ZM, Sebesta JA and Brown TA (2008) Primary tumor location impacts breast cancer survival Am J Surg 195(5) 641–644 DOI: 10.1016/j.amjsurg.2007.12.039 PMID: 18424280

12. Lohrisch C, Jackson J, Jones A, Mates D and Olivotto IA (2000) Relationship between tumor location and relapse in 6,781 women with early invasive breast cancer J Clin Oncol 18(15) 2828–2835 PMID: 10920130

13. Sarp S, Fioretta G, Verkooijen HM and Vlastos G (2007) Tumor location of the lower-inner quadrant is associated with an impaired survival for women with early-stage breast cancer Ann Surg Oncol 14(3) 1031–1039 DOI: 10.1245/s10434-006-9231-5

14. Wu S, Zhou J, Ren Y, Sun J and Li F et al (2014) Tumor location is a prognostic factor for survival of Chinese women with T1-2N0M0 breast cancer Int J Surg 12(5) 394–398 DOI: 10.1016/j.ijsu.2014.03.011 PMID: 24657350

15. Brautigam E, Track C, Seewald DH and Feichtinger J et al (2009) Medial tumor localization in breast cancer—an unappreciated risk factor? Strahlenther Onkol 185(10) 663–668 DOI: 10.1007/s00066-009-1984-x PMID: 19806331

16. Hazrah P, Dhir M, Gupta SD, Deo V and Parshad R (2009) Prognostic significance of location of the primary tumor in operable breast cancers Indian J Cancer 46(2) 139–145 DOI: 10.4103/0019-509X.49152 PMID: 19346648

17. Kamakura T, Akazawa K, Nomura Y and Sugimachi K et al (1996) Poor prognosis of lower quadrant breast carcinoma. Nishi Nippon study group on adjuvant chemo-endocrine therapy for breast cancer J Surg Oncol 61(4) 295–299 PMID: 8628002

18. Boyages J, Recht A and Connolly JL et al (1990) Early breast cancer: predictors of breast recurrence for patients treated with conservative surgery and radiation therapy Radiother Oncol 19(1) 29–41 DOI: 10.1016/0167-8140(90)90163-Q PMID: 2173044

19. Fowble B, Solin LJ, Schultz DJ and Weiss MC (1992) Breast recurrence and survival related to primary tumor location in patients undergoing conservative surgery and radiation for early-stage breast cancer Int J Radiat Oncol Biol Phys 23(5) 933–939 DOI: 10.1016/0360-3016(92)90897-Q PMID: 1322386

20. Fisher B, Wolmark N, Redmond C, Deutsch M and Fisher ER (1981) Findings from NSABP protocol no. B-04: comparison of radical mastectomy with alternative treatments II The clinical and biologic significance of medial-central breast cancers Cancer 48(8) 1863–1872 DOI: 10.1002/1097-0142(19811015)48:8<1863::AID-CNCR2820480825>3.0.CO;2-U PMID: 7284980

21. Veronesi U, Marubini E, Marianni L, Valagussa P and Zucali R (1999) The dissection of internal mammary nodes does not improve the survival of breast cancer patients 30-year results of a randomised trial Eur J Cancer 35(9) 1320–1325 DOI: 10.1016/S0959-8049(99)00133-1

22. Hennequin C, Bossard N and Servag-Vernat S et al (2013) Ten-year survival results of a randomized trial of irradiation of internal mammary nodes after mastectomy Int J Radiat Oncol Biol Phys 86(5) 860–866 DOI: 10.1016/j.ijrobp.2013.03.021 PMID: 23664327

23. Fowble B, Hanlon A, Freedman G and Nicolaou N et al (2000) Internal mammary node irradiation neither decreases distant metastases nor improves survival in stage I and II breast cancer Int J Radiat Oncol Biol Phys 47(4) 883–894 DOI: 10.1016/S0360-3016(00)00526-5 PMID: 10863056
24. Ellsworth RE, Zhu K, Bronfman L and Gutchell V et al (2008) The clinical breast care project: an important resource in investigating environmental and genetic contributions to breast cancer in African American women Cell Tissue Bank 9(2) 109–120 DOI: 10.1007/s10561-007-9054-z

25. American Joint Committee on C (2010) AJCC Cancer Staging Manual, vol 7 (New York: Springer)

26. Elston CW and Ellis IO (1991) Pathological prognostic factors in breast cancer. I. The value of histological grade in breast cancer: experience from a large study with long-term follow-up Histopathology 19(5) 403–410 DOI: 10.1111/j.1365-2559.1991.tb00229.x PMID: 1757079

27. Bloom HJ and Richardson WW (1957) Histological grading and prognosis in breast cancer; a study of 1409 cases of which 359 have been followed for 15 years Br J Cancer 11(3) 359–377 DOI: 10.1038/bjc.1957.43 PMID: 13499785 PMCID: 2073885

28. Hammond MEH, Hayes DF and Dowsett M et al (2010) American society of clinical oncology/college of American pathologists guideline recommendations for immunohistochemical testing of estrogen and progesterone receptors in breast cancer J Clin Oncol 28(16) 2784–2795 DOI: 10.1200/JCO.2009.25.6529 PMID: 20404251 PMCID: 2881855

29. Carter CL, Allen C and Henson DE (1989) Relation of tumor size, lymph node status, and survival in 24,740 breast cancer cases Cancer 63(1) 181–187 DOI: 10.1002/1097-0142(19890101)63:1<181::AID-CNCR2820630129>3.0.CO;2-H PMID: 2910416

30. Darbre PD (2005) Recorded quadrant incidence of female breast cancer in Great Britain suggests a disproportionate increase in the upper outer quadrant of the breast Anticancer Res 25(3c) 2543–2550 PMID: 16080490

31. Aljarrah A and Miller W (2014) Trends in the distribution of breast cancer over time in the southeast of Scotland and review of the literature Ecancermedicalscience 8 427 PMID: 24834122 PMCID: 4019459

32. Smeets A, Ryckx A, Belmans A and Wildiers H et al (2013) Impact of tumor chronology and tumor biology on lymph node metastasis in breast cancer Springer Plus 2(1) 480 DOI: 10.1186/2193-1801-2-480 PMID: 24083118 PMCID: 3786086

33. Laura S, Coombs NJ, Ung O and Boyages J (2006) Tumour size as a predictor of axillary node metastases in patients with breast cancer ANZ J Surg 76(11) 1002–1006 DOI: 10.1111/j.1445-2197.2006.03918.x PMID: 17054550

34. Yoshihara E, Smeets A and Laenen A et al (2013) Predictors of axillary lymph node metastases in early breast cancer and their applicability in clinical practice Breast 22(3) 357–361 DOI: 10.1016/j.breast.2012.09.003

35. Postaci H, Zengel B, Yararbas U and Uslu A et al (2013) Sentinel lymph node biopsy in breast cancer: predictors of axillary and non-sentinel lymph node involvement Balkan Med J 30(4) 415–421 DOI: 10.5152/balkanmedi.2013.9591

36. Wu JL, Tseng HS, Yang LH and Wu HK et al (2014) Prediction of axillary lymph node metastases in breast cancer patients based on pathologic information of the primary tumor Med Sci Monit 20 577–581 DOI: 10.12659/MSM.890345 PMID: 24714517 PMCID: 3989944

37. Greer LT, Rosman M, Charles Mylander W and Liang W et al (2014) A prediction model for the presence of axillary lymph node involvement in women with invasive breast cancer: a focus on older women Breast J 20(2) 147–153 DOI: 10.1111/tbj.12233 PMID: 24475876

38. Friedman D, Gipponi M and Murelli F et al (2013) Predictive factors of non-sentinel lymph node involvement in patients with invasive breast cancer and sentinel node micrometastases Anticancer Res 33(10) 4509–4514 PMID: 24123023

39. Mojarad S, Venturini B and Fulgenzi P et al (2013) Prediction of nodal metastasis and prognosis of breast cancer by ANN-based assessment of tumour size and p53, Ki-67 and steroid receptor expression Anticancer Res 33(9) 3925–3933 PMID: 24023330

40. Gurleyik G, Aker F, Aktekin A and Saglam A (2011) Tumor characteristics influencing non-sentinel lymph node involvement in clinically node negative patients with breast cancer J Breast Cancer 14(2) 124–128 DOI: 10.4048/jbc.2011.14.2.124 PMID: 21847407 PMCID: 3148535
41. Aitken E and Osman M (2010) Factors affecting nodal status in invasive breast cancer: a retrospective analysis of 623 patients. *Breast J* **16**(3) 271–278 DOI: 10.1111/j.1524-4741.2009.00897.x PMID: 20210804

42. Michaelson JS, Silverstein M, Wyatt J and Weber G et al (2002) Predicting the survival of patients with breast carcinoma using tumor size. *Cancer* **95**(4) 713–723 DOI: 10.1002/cncr.10742 PMID: 12209713

43. Narod SA (2012) Tumour size predicts long-term survival among women with lymph node-positive breast cancer. *Curr Oncol* **19**(5) 249–253 DOI: 10.3747/co.19.1043 PMID: 23144572 PMCID: 3457875

44. Klintman M, Nilsson F, Bendahl PO and Ferno M et al (2013) A prospective, multicenter validation study of a prognostic index composed of S-phase fraction, progesterone receptor status, and tumour size predicts survival in node-negative breast cancer patients: NNBC, the node-negative breast cancer trial. *Ann Oncol* **24**(9) 2284–2291 DOI: 10.1093/annonc/mdt186 PMID: 23704202

45. Gasparini G, Weidner N, Bevilacqua P and Maluta S et al (1994) Tumor microvessel density, p53 expression, tumor size, and peritumoral lymphatic vessel invasion are relevant prognostic markers in node-negative breast carcinoma. *J Clin Oncol* **12**(3) 454–466 PMID: 7509851

46. Michaelson JS, Silverstein M, Sgroi D and Cheongsiatmoy JA et al. (2003) The effect of tumor size and lymph node status on breast carcinoma lethality. *Cancer* **98**(10) 2133–2143 DOI: 10.1002/cncr.11765 PMID: 14601082

47. Nicholson BT, Harvey JA and Cohen MA (2009) Nipple-areolar complex: normal anatomy and benign and malignant processes. *Radiographics* **29**(2) 509–523 DOI: 10.1148/rg.292085128 PMID: 19325062

48. Giess CS, Keating DM, Osborne MP, Ng YY and Rosenblatt R (1998) Retroareolar breast carcinoma: clinical, imaging, and histo-pathologic features. *Radiology* **207**(3) 669–673 DOI: 10.1148/radiology.207.3.9609889 PMID: 9609889

49. Feig SA, Shaber GS, Patchefsky A and Schwartz GF et al (1977) Analysis of clinically occult and mammographically occult breast tumors. *AJR Am J Roentgenol* **128**(3) 403–408 DOI: 10.2214/ajr.128.3.403 PMID: 190908

50. Da CD, Taddese A and Cure ML et al (2007) Common and unusual diseases of the nipple-areolar complex. *Radiographics* **27** Suppl 1 S65–S77