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Tumor location of the central and nipple portion is associated with impaired survival for women with breast cancer

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Background: Tumor location in the breast varies, with the highest frequency in the upper outer quadrant and lowest frequency in the lower inner quadrant. Nevertheless, tumors in the central and nipple portion (TCNP) are poorly studied types of breast cancer; therefore, we aimed to clarify the clinicopathological characteristics and prognostic features of TCNP.

Methods: Using the Surveillance, Epidemiology, and End Results database, we identifed 105,037 patients diagnosed with tumor in the breast peripheral quadrant (TBPQ) (n=97,046) or TCNP (n=7,991). The chi-squared test was used to compare categorical variables across TCNP and TBPQ. Cox proportional hazard models with hazard ratios were applied to estimate the factors associated with prognosis.

Results: The median follow-up was over 43 months. Compared with TBPQ, TCNP patients were significantly older (age ≥66 years: 40.4% vs 34.1%, P<0.001), with larger tumor sizes (>20 mm size: 46.9% vs 37.3%, P<0.001), higher proportions of TNM stage II-III (18.6% vs 9.9%, P<0.001), and more mastectomies (58.1% vs 37.8%, P<0.001). The breast cancer-specifc survival (BCSS)/overall survival (OS) rate was significantly worse for TCNP than for TBPQ. Multivariate Cox analysis showed a higher hazard ratios for TCNP over TBPQ (BCSS: hazard ratios =1.160, P=0.005, 95% CI: 1.046-1.287; OS: hazard ratios =1.301, P<0.001, 95% CI: 1.211-1.398). A subgroup analysis revealed inferior outcomes for TCNP in TNM stage II-III and breast subtype subgroup. Multivariate logistic regression indicated that TCNP was an independent contributing factor to LN metastasis.

Conclusions: TCNP was associated with older age, larger tumor size, higher TNM stage, and lymph node metastasis. Compared with TBPQ, TCNP had adverse impacts on BCSS and OS. **Keywords:** tumor in the central and nipple portion, tumor in the breast peripheral quadrant, prognosis, lymph node metastasis

Introduction

Breast cancer is the most common malignancy in women and one of the three most common cancers worldwide. Although mortality from breast cancer in North America and the European Union has decreased, breast cancer remains the most common cause of death from cancer in less-developed countries and is second to lung cancer in more developed countries. 1,2 Therefore, it is essential to find effective parameters to identify patients with poor survival and to support them with personalized therapy.

A number of pathological characteristics, including tumor size and grade, hormone receptor (HR) status, human epidermal growth factor receptor-2 (HER2), and lymph node (LN) status, have been identified, all of which are considered when predicting prognosis and determining the most effective treatment options.^{3–5} Furthermore, molecular tests have been developed that add prognostic value by determining intrinsic subtype, stratification of tumors into low or high grade, or predicting recurrence. 6-9 Despite these factors and tests, predicting patient outcome is not truly enough, as patients with similar pathological characteristics treated with identical regimens often exhibit highly variable clinical outcomes, indicating that additional prognostic factors are still need to be identified to improve patient stratification.

Tumor location in the breast has been reported as an independent prognostic factor. Tumors in the upper outer quadrant (UOQ) are the most frequent site of tumor location. They are associated with better survival compared with other quadrants; 10,11 survival for non-UOQ tumors such as lower inner quadrant (LIQ) or medial regions have demonstrated lower survival. 12-17 Axillary LN metastasis has also been suggested to be significantly lower in the upper inner quadrant (UIQ, 20.6%) compared with all other quadrants (33.2%).18 Nevertheless, studies regarding tumor in the central and nipple portion (TCNP) of the breast are discordant. One study suggested that breast cancer located closer to the nipple had a higher incidence of metastases to axillary LNs, ¹⁹ while another demonstrated that there was no evidence that correlated intramammary tumor proximity to the nipple with the presence of axillary LN metastasis.²⁰ Furthermore, the clinicopathological characteristics and the effect of TCNP on survival have still not been deeply explored.

Therefore, the aim of this study was to investigate the clinicopathological characteristics of TCNP and to clarify their prognostic value using a population-based database.

Materials and methods Database

We used the Surveillance, Epidemiology, and End Results (SEER) database (November 2017 submission), a National Cancer Institute-sponsored program, and obtained data from 18 population-based registries with the SEER*Stat software, version 8.3.4 (http://seer.cancer.gov/about/).

Study population

We derived a dataset of female breast cancer patients diagnosed from 2010 to 2013, for the reason of known HER2 status and follow-up time. The inclusion criteria were as follows: females aged over 18 years old; American Joint Committee on Cancer stages (the seventh AJCC System) TNM stage I–III;^{21,22} unilateral breast cancer; breast cancer as the first and only cancer diagnosis; TCNP or tumor in the breast peripheral quadrant (TBPQ) (including UOQ, UIQ, lower outer quadrant, LIQ, but overlapping sites were

excepted); diagnosis not obtained from a death certificate or autopsy; pathologic confirmation of breast cancer; known tumor differentiation status (histological score);^{23–25} known HR and HER2 status;^{26–28} known tumor size and LN status; known surgery type; known radiation/chemotherapy; and active follow-up. Patients with incomplete information were excluded. In total, 105,037 patients were included, including 7,991 TCNP (7.60%) and 97,046 TBPQ patients (92.40%).

Ethics statement

This study was approved by an independent ethical institutional review board at Guangdong Provincial People's Hospital and was carried out in accordance with the principles of the Declaration of Helsinki. The data released by the SEER database were publicly available and do not require patient's informed consent.

Statistical analysis

Statistical analysis was performed using SPSS software for Windows version 25.0 (SPSS Inc., Chicago, IL, USA). The chi-squared test was used to compare categorical variables across TCNP and TBPQ. Breast cancer-specific survival (BCSS) was defined as the survival time from the diagnosis date of breast cancer to the date of death caused by breast cancer, and the overall survival (OS) was defined as the time from the diagnosis date of breast cancer to the date of death from any cause. Kaplan-Meier curves, competing risks analysis (for BCSS), and log-rank tests (for OS) were generated to determine differences in the survival analyses. In addition, Cox proportional hazard models with hazard ratios were applied to estimate the factors associated with prognosis. A 1:1 paired match, by the propensity score matching (PSM) method, was also carried out to balance the impact of the baseline. Logistic regression was utilized to present the relationship between the tumor location and axillary LN metastasis. Two-sided P<0.05 was considered statistically significant, and reported P-values were not corrected for multiple testing.

Results

Demographic and clinical characteristics of TCNP

We first summarized the demographic and clinical characteristics of all 105,037 patients (Table 1). The median follow-up was over 43 months. Compared with the TBPQ patients, TCNP patients were significantly older (age \geq 66 years: 40.4% vs 34.1%, P<0.001); they had a larger tumor size (>20 mm size: 46.9% vs 37.3%, P<0.001) and higher TNM stage II–III (18.6% vs 9.9%, P<0.001). More TCNP

Table I Baseline characteristics of patients with TCNP or TBPQ from the SEER database 2010–2013

Variables	Data before PSM		P-value	Data after PSM		P-value
	TCNP TBPQ			TCNP TBPQ		
	(7,991)	(97,046)		(7,991)	(7,991)	
Age (years), %			<0.001			0.945
18–45	11.0	14.0		11.0	11.0	
46–55	21.3	24.4		21.3	21.1	
56–65	27.3	27.5		27.3	27.8	
66–79	28.5	26.5		28.5	28.5	
>80	11.9	7.6		11.9	11.6	
Race, %			<0.001			0.087
White	79.9	79.5		79.9	81.2	
Black	9.2	10.8		9.2	9.1	
Others	10.9	9.7		10.9	9.7	
Marital status, %			0.744			0.929
Married	81.3	81.0		81.5	81.3	
Not married	18.7	19.0		18.5	18.7	
Laterality, %			0.576			0.962
Left	51.3	50.9		51.3	51.3	
Right	48.7	49.1		48.7	48.7	
Histology, %			<0.001			0.617
IDC	70.8	77.1		70.8	71.3	
ILC	10.4	8.4		10.4	10.5	
Others	18.8	14.5		18.8	18.2	
Tumor differentiation, %			<0.001			0.103
I–II	67.1	65.3		67.I	67.3	
III–IV	32.9	34.7		32.9	32.7	
TNM stage, %			<0.001			0.861
	81.4	90.1		43.7	43.9	
II–III	18.6	9.9		56.3	56.1	
Tumor size (mm), %			<0.001			0.111
0–20	53.1	62.7	10000	54.1	53.5	
21–50	35.1	31.1		35.3	36.6	
>50	11.8	6.2		10.6	9.9	
LN status, %			<0.001			0.871
Negative	61.3	72.2	10.001	61.3	61.1	
Positive	38.7	27.8		38.7	38.9	
Breast subtype, %			<0.001	155	33.7	0.174
HER2-/HR+	76.0	74.0	~0.001	76.0	77.1	
HER2+/HR+	10.8	9.7		10.8	10.1	
HER2+/HR-	4.9	4.0		4.9	4.4	
Triple-negative	8.3	12.3		8.3	8.4	
Surgery, %	0.3	12.3	<0.001	0.3	0.7	0.873
	41.0	(2.2	<0.001	41.0	41.0	0.073
Lumpectomy	41.9	62.2		41.9	41.8	
Mastectomy	58.1	37.8	-0.001	58.1	58.2	0.410
Radiation, %	47.2	F(0	<0.001	47.3	47.5	0.419
Yes	47.3	56.9		47.3	47.5	
No/unknown	52.7	43.1	0.222	52.7	52.5	0.007
Chemotherapy, %	42.2	41.5	0.233	42.2	42.2	0.987
Yes	42.2	41.5		42.2	42.2	
No/unknown	57.8	58.5		57.8	57.8	

Abbreviations: HER2, human epidermal growth factor receptor-2; HR, hormone receptor; IDC, invasive ductal carcinoma; ILC, invasive lobular carcinoma; LN, lymph node; PSM, propensity score matching; SEER, Surveillance, Epidemiology, and End Results; TBPQ, tumor in the breast peripheral quadrant; TCNP, tumor in the central and nipple portion.

patients underwent mastectomy (58.1% vs 37.8%, *P*<0.001). Furthermore, TCNP had a significantly higher LN positivity rate (LN+: 38.7% vs 27.8%, *P*<0.001). These characteristics suggested that TCNP had a clinicopathological baseline distinct from that of TBPQ.

Comparison of survival between TCNP and TBPO

To evaluate the BCSS and OS of TCNP and TBPQ, the Kaplan–Meier method was used. The 5-year BCSS rates of the patients with TCNP were significantly lower than those of the TBPQ population (92.1% vs 94.9%, *P*<0.001) (Figure 1A). Furthermore, the 5-year OS rates of the patients with TCNP were markedly lower than those of the TBPQ population (84.8% vs 92.5%, *P*<0.001) (Figure 1B).

Because of the distinct clinicopathological baseline of TCNP, univariate and multivariate Cox proportional hazards models were used to balance the effect of factors.

Age, race, marital status, location of TCNP or TBPQ, tumor differentiation, TNM stage, tumor size, LN status, breast subtype, surgery type, radiation, and chemotherapy (all P<0.001) were responsible for BCSS. Similarly, the significant predictors of OS in patients were race, marital status, location of TCNP or TBPQ, TNM stage, tumor size, LN status, surgery type, and radiation (all P<0.001) (Table 2). After multivariate analysis, we found that race, marital status, location of TCNP or TBPQ, tumor differentiation, TNM stage, tumor size, LN status, breast subtype, surgery type, and radiation were significant independent predictors of BCSS (all P<0.05), while race, location of TCNP or TBPQ, tumor differentiation, TNM stage, tumor

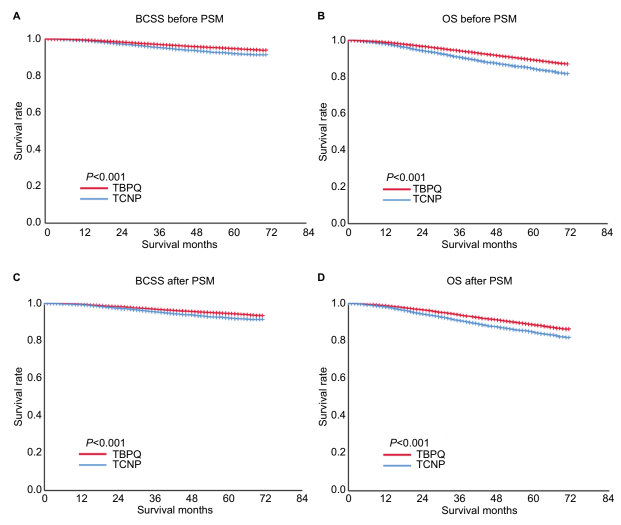


Figure I Relationship between TCNP/TBPQ and BCSS/OS of breast cancer patients.

Notes: (A) BCSS of patients with TCNP was significantly shorter than those with TBPQ, including BCSS after PSM (C). (B) OS of patients with TCNP was also markedly shorter than those with TBPQ, including BCSS after PSM (D).

Abbreviations: BCSS, breast cancer specific survival; OS, overall survival; PSM, propensity score matching; SEER, Surveillance, Epidemiology, and End Results; TBPQ, tumor in the breast peripheral quadrant; TCNP, tumor in the central and nipple portion.

Table 2 Univariate Cox regression model analysis of BCSS and OS between TCNP and TBPQ

Variables	BCSS		os					
	Hazard ratios	95% CI	P-value	Hazard ratios	95% CI	P-value		
Age (years)								
18–45	Reference			Reference				
46–55	0.729	0.660-0.805	<0.001	0.818	0.748-0.894	<0.001		
56–65	0.660	0.598-0.729	<0.001	0.971	0.892-1.056	0.490		
66–79	0.732	0.664-0.808	<0.001	1.640	1.515–1.774	<0.001		
≥80	1.685	1.511–1.879	<0.001	5.433	5.017–5.884	<0.001		
Race	I							
White	Reference			Reference				
Black	1.908	1.763-2.064	<0.001	1.471	1.386-1.560	<0.001		
Others	0.690	0.606–0.787	<0.001	0.584	0.530-0.642	<0.001		
Marital status			10000			101001		
Married	Reference			Reference				
Unmarried	1.392	1.283-1.509	<0.001	1.122	1.056-1.192	<0.001		
Location		1			1	10.001		
TBPQ	Reference			Reference	I			
TCNP	1.532	1.384–1.696	<0.001	1.522	1.418-1.634	<0.001		
Laterality	1	1	1	1	1	1		
Left	Reference			Reference				
Right	0.946	0.889-1.005	0.074	0.971	0.931-1.013	0.971		
Histology								
IDC	Reference			Reference				
ILC	0.743	0.657-0.840	<0.001	0.960	0.889-1.073	0.303		
Others	0.822	0.749-0.901	<0.001	0.978	0.921-1.039	0.472		
Tumor different			10.00	1	11121 11121			
I–II	Reference				Reference			
III–IV	4.687	4.376–5.020	<0.001	2.155	2.062–2.253	<0.001		
TNM stage								
l	Reference			Reference				
-	7.225	6.625–7.879	<0.001	2.719	2.597–2.846	<0.001		
Tumor size (mm	1)							
0–20	Reference			Reference				
21–50	4.209	3.909-4.532	<0.001	2.361	2.255–2.472	<0.001		
>50	10.965	10.032-11.983	<0.001	4.540	4.253-4.874	<0.001		
LN status	1	1	10.001	1 17 1	1	1 10.001		
Negative	Reference			Reference				
Positive	4.435	4.160–4.729	<0.001	2.139	2.050–2.232	<0.001		
Breast subtype		1	1	1	1 ,	1 10.001		
HER2-/HR+	Reference			Reference				
HER2+/HR+	1.343	1.199–1.505	<0.001	1.018	0.941-1.101	0.656		
HER2+/HR-	2.818	2.496–3.181	<0.001	1.606	1.460–1.767	<0.001		
Triple-negative	4.810	4.495–5.147	<0.001	2.557	2.430–2.690	<0.001		
Surgery	1	1 3 17	\0.001	1		\0.001		
Lumpectomy	Reference			Reference				
Mastectomy	2.671	2.507–2.846		1.759	1.686-1.836	<0.001		
Radiation	1 1	2.557 2.510		1 = 7	1	\0.001		
Yes	Reference			Reference				
No	1.505	1.413–1.605	<0.001	1.861	1.780–1.946	<0.001		
Chemotherapy	1.505	1.113 1.003	\0.001	1.001	1.700-1.740	\<0.001		
Yes	Reference			Reference				
No/unknown	0.389	0.364-0.416	<0.001	0.984	0.941-1.029	0.483		
1 40/ UTINITO WIT	0.507	0.301 0.410	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	0.701	0.711-1.027	0. 103		

Abbreviations: BCSS, breast cancer specific survival; HER2, human epidermal growth factor receptor-2; HR, hormone receptor; IDC, invasive ductal carcinoma; ILC, invasive lobular carcinoma; LN, lymph node; OS, overall survival; TBPQ, tumor in the breast peripheral quadrant; TCNP, tumor in the central and nipple portion.

size, LN status, surgery type, and radiation were significant independent predictors of OS (all P<0.05). The tumor location of TCNP was confirmed as an independent risk factor over TBPQ in terms of both BCSS and OS (BCSS: hazard ratios =1.160, P=0.005, 95% CI: 1.046–1.287; OS: hazard ratios =1.301, P<0.001, 95% CI: 1.211–1.398) (Table 3).

Survival analysis in the matched group

A 1:1 (TCNP/TBPQ) matched case-control analysis by PSM was performed to balance the baseline differences. In

total, 7,991 patients were obtained for each location type. Ten factors were included in the PSM, including age, race, marital status, laterality, histology, tumor differentiation, TNM stage, tumor size, LN status, breast subtype, surgery type, radiation, and chemotherapy. No statistically significant differences were found in terms of the characteristics between the two groups (Table 1). The survival analysis of the matched groups, which was consistent with the entire cohort analysis, showed that TCNP presented a worse outcome for the BCSS (5-year BCSS: 92.1% vs 94.5%, *P*<0.001) (Figure 1C) and a

Table 3 Multivariate Cox regression model analysis of the BCSS and OS between TCNP and TBPQ

Variables	BCSS			os				
	Hazard ratios	95% CI	P-value	Hazard ratios	95% CI	P-value		
Age (years)			·		•			
18–45	Reference							
46–55	1.045	0.942-1.160	0.401					
56–65	1.173	1.056-1.302	0.003					
66–79	1.609	1.449-1.787	<0.001					
≥80	3.521	3.130–3.960	<0.001					
Race	·	· ·		'	'			
White	Reference			Reference				
Black	1.260	1.158-1.372	<0.001	1.207	1.134–1.285	<0.001		
Others	0.686	0.598-0.787	<0.001	0.543	0.491-0.600	<0.001		
Marital status		I						
Married	Reference		,					
Unmarried	1.203	1.106-1.308	<0.001					
Location		I						
TBPQ	Reference		,	Reference				
TCNP	1.160	1.046-1.287	0.005	1.301	1.211-1.398	<0.001		
Tumor differentiation	1	,		,				
I–II	Reference			Reference				
III–IV	2.412	2.228-2.612	<0.001	1.652	1.578-1.730	<0.001		
TNM stage		,		,	'			
I	Reference			Reference				
II–III	1.826	1.579-2.112	<0.001	1.109	1.004-1.224	0.041		
Tumor size (mm)		,		,	'			
0–20	Reference			Reference				
21–50	1.535	1.375-1.714	<0.001	1.643	1.507-1.791	<0.001		
>50	3.187	2.818-3.604	<0.001	3.116	2.823-3.439	<0.001		
LN status	1							
Negative	Reference			Reference				
Positive	2.474	2.283–2.681	<0.001	1.509	1.427–1.595	<0.001		
Breast subtype				I				
HER2-/HR+	Reference	,	,					
HER2+/HR+	0.846	0.750-0.955	0.007					
HER2+/HR-	1.358	1.192–1.547	<0.001					
Triple-negative	2.590	2.391–2.806	<0.001					
Surgery			10.001		1			
Lumpectomy	Reference	Reference			Reference			
Mastectomy	1.271	1.179–1.370	<0.001	0.905	0.859-0.952	<0.001		
Radiation			10.001	1 *** * *	1	1 .3.001		
Yes	Reference			Reference				
	1			1				

Abbreviations: BCSS, breast cancer specific survival; HER2, human epidermal growth factor receptor-2; HR, hormone receptor; LN, lymph node; OS, overall survival; TBPQ, tumor in the breast peripheral quadrant; TCNP, tumor in the central and nipple portion.

statistically worse outcome for the OS (5-year OS: 84.3% vs 92.1%, *P*<0.001) compared with TBPQ (Figure 1D).

Survival analysis in various subgroups

The analyses above verified that TCNP was an independent prognostic factor and significantly negatively correlated with BCSS and OS. We next assessed the prognostic value of TCNP compared with TBPQ in various subgroups. We found that TCNP was a worse prognostic indicator for

BCSS (*P*<0.001) and OS (*P*<0.001) in patients with TNM stage II–III (Figure 2A, B), and this poor prognostic value of BCSS and OS of TCNP also existed for patients with HER2⁻/HR⁺ (BCSS: *P*=0.001; OS: *P*<0.001) (Figure 3A, E), HER2⁺/HR⁺ (BCCS: *P*<0.001; OS: *P*<0.001) (Figure 3C, G), and triple-negative cancer (Figure 3D, H) (BCSS: *P*<0.001; OS: *P*<0.001). Despite the fact that prognostic value of BCSS of TCNP in patients with HER2⁺/HR⁻ showed no significance (*P*=0.538), OS of TCNP in patients

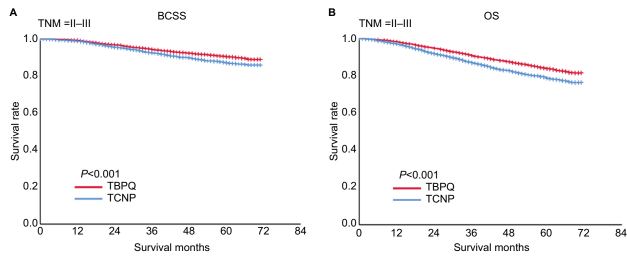


Figure 2 Kaplan–Meier survival curves of different breast cancer subgroups.

Note: TCNP significantly correlated with shorter BCSS and OS in subgroups with TNM stage II–III (A, B).

Abbreviations: BCSS, breast cancer specific survival; OS, overall survival; TBPQ, tumor in the breast peripheral quadrant; TCNP, tumor in the central and nipple portion.

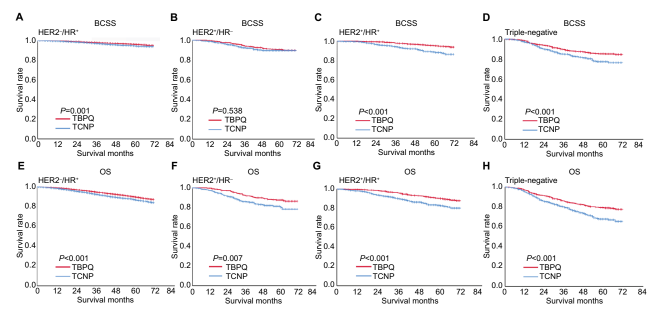


Figure 3 Kaplan–Meier survival curves of different breast cancer subgroups.

Note: TCNP significantly correlated with shorter BCSS and OS in subgroups with various breast subtypes (A–H).

Abbreviations: BCSS, breast cancer specific survival; HER2, human epidermal growth factor receptor-2; HR, hormone receptor; OS, overall survival; TBPQ, tumor in the breast peripheral quadrant: TCNP, tumor in the central and nipple portion.

with HER2 $^+$ /HR $^-$ was significant (P=0.007) (Figure 3B, F and Table 4).

TCNP related to LN metastases of breast cancer

To clarify the reasons leading to poor prognosis of TCNP, we analyzed the relationship between TCNP and LN metastases of breast cancer. Variables that were significant (P<0.001) in the univariate logistic analysis (age, race, tumor location, histology, tumor differentiation, TNM stage, tumor size, and breast subtype) were further included in the multivariate logistic regression analysis. TCNP was more likely to metastasize to LNs (OR =3.903, 95% CI: 3.687–4.133, P<0.001) (Table 5). This result verified the linkage of TCNP and LN metastasis and may partially explain the worse outcome for TCNP.

Discussion

Determination of whether tumor location can be used prognostically is important in optimizing treatment. TCNP has a unique anatomical position, but its clinicopathological and survival characteristics are not well investigated. In this study, we showed that TCNP, compared with TBPQ, has distinct clinicopathological characteristics and worse survival.

First, we found that TCNP patients were mainly in an older age group (age ≥66 years), with larger tumor sizes (>20 mm size) and higher TNM stages (II–III). This may because tumors within the central region are harder to detect for reasons of overpenetration of X-rays in the nipple–areolar

complex and may reach a substantially larger size before being detected by imaging;²⁹ therefore, TCNP relates to tumor burden and progression. By univariate analysis, we found many significant prognostic factors for BCSS or OS, including age, race, marital status, location of TCNP or TBPQ, tumor differentiation, TNM stage, tumor size, LN status, breast subtype, surgery type, radiation, and chemotherapy. After multivariate analysis, however, we discovered that only race, location of TCNP or TBPQ, tumor differentiation, TNM stage, tumor size, LN status, surgery type, and radiation were significant independent predictors of BCSS and OS. Tumor location of TCNP was finally confirmed as an independent risk factor over TBPQ for both BCSS and OS.

Through further analysis, we found that shorter survival of TCNP patients was detected in this study. The 1-, 3-, and 5-year BCSS rates and OS rates of patients with TCNP were markedly lower than those of patients with TBPQ, including the survival analysis after PSM. The mechanisms may be the understaging of TCNP at diagnosis. Ansari et al¹⁹ reported that breast cancers located closer to the nipple have a higher incidence of metastases to axillary LN s. In our study, TCNP had a significantly higher LN positivity rate than TBPQ (LN+: 38.7% vs 27.8%, P<0.001). We also conducted a multivariate logistic regression analysis that indicated that TCNP was an independent indicator for LN metastasis, and metastasis is the fundamental cause of higher mortality and poorer prognosis of patients who suffer from malignant tumors. Furthermore, the difficulty of mammographic detection and more frequent occurrence of

Table 4 BCSS rates, OS rates, and P-values for various subgroups

Groups	BCSS				os					
	I-year	3-year	5-year	P-value	I-year	3-year	5-year	P-value		
TNM II-III										
TBPQ	99.1%	94.2%	90.3%	<0.001	98.1%	90.7%	83.8%	<0.001		
TCNP	98.7%	92.4%	87.0%		97.0%	86.9%	78.6%			
HER2-/HR+										
TBPQ	99.6%	97.8%	95.9%	0.001	98.8%	94.7%	89.6%	<0.001		
TCNP	99.5%	96.9%	94.4%		98.3%	92.3%	86.7%			
HER2+/HR+										
TBPQ	100%	97.8%	95.2%	<0.001	99.6%	95.6%	90.2%	<0.001		
TCNP	99.4%	94.2%	88.3%		97.9%	89.8%	82.6%			
HER2+/HR-										
TBPQ	99.0%	93.8%	90.4%	0.538	99.7%	91.9%	87.6%	0.007		
TCNP	98.4%	92.2%	89.4%		97.2%	85.6%	79.8%			
Triple-negativ	re e									
TBPQ	98.1%	89.4%	85.2%	<0.001	96.7%	85.4%	79.0%	<0.001		
TCNP	96.9%	84.8%	77.3%		94.8%	79.5%	67.6%			

Abbreviations: BCSS, breast cancer specific survival; HER2, human epidermal growth factor receptor-2; HR, hormone receptor; OS, overall survival; TBPQ, tumor in the breast peripheral quadrant; TCNP, tumor in the central and nipple portion.

Table 5 Univariate and multivariate logistic regression analysis on LN metastasis

Variables	Univaria	e analysis		Multivariate analysis				
	OR	95% CI	P-value	OR	95% CI	P-value		
Age (years)	'		'	'		,		
18–45	Reference			Reference				
46–55	0.486	0.465-0.507	<0.001	0.852	0.805-0.902	<0.001		
56–65	0.360	0.345-0.375	<0.001	0.764	0.722-0.809	<0.001		
66–79	0.290	0.278-0.303	<0.001	0.658	0.620-0.698	<0.001		
≥80	0.898	0.848-0.951	<0.001	0.733	0.679-0.793	<0.001		
Race	'		'		<u>'</u>	· ·		
White	Reference			Reference	Reference			
Black	3.364	3.245–3.487	<0.001	1.585	1.500-1.675	<0.001		
Others	2.808	2.708–2.912	<0.001	1.571	1.484–1.663	<0.001		
Location								
TBPQ	Reference	Reference			Reference			
TCNP	5.247	5.106–5.393	<0.001	3.903	3.687–4.133	<0.001		
Histology	·							
IDC	Reference			Reference				
ILC	4.612	4.446–4.784	<0.001	1.976	1.861-2.098	<0.001		
Others	1.772	1.713–1.813	<0.001	1.164	1.106–1.225	<0.001		
Tumor differentiation			•					
I–II	Reference			Reference				
III–IV	2.905	2.831-2.982	<0.001	1.145	1.097-1.196	<0.001		
TNM stage			•			•		
I	Reference			Reference				
II–III	12.086	11.726–12.457	<0.001	191.671	175.214–209.674	<0.001		
Tumor size (mm)								
0–20	Reference	Reference			Reference			
21–50	3.882	3.767-4.001	<0.001	0.057	0.052-0.062	<0.001		
>50	18.943	18.150–19.770	<0.001	0.207	0.187-0.229	<0.001		
Breast subtype								
HER2-/HR+	Reference			Reference				
HER2+/HR+	0.376	0.369–0.382	<0.001	1.517	1.432-1.606	<0.001		
HER2+/HR-	0.539	0.517-0.563	<0.001	2.688	2.479–2.915	<0.001		
Triple-negative	0.625	0.586-0.667	<0.001	0.852	0.805-0.902	<0.001		

Abbreviations: BCSS, breast cancer specific survival; HER2, human epidermal growth factor receptor-2; HR, hormone receptor; IDC, invasive ductal carcinoma; ILC, invasive lobular carcinoma; LN, lymph node; OS, overall survival; TBPQ, tumor in the breast peripheral quadrant; TCNP, tumor in the central and nipple portion.

palpable tumors in the central region not only explained the larger tumor size but also explained the increased mortality found in our study.

We elucidated the prognostic value of TCNP in various subgroups of breast cancer patients. Compared with TBPQ, we found that TCNP gave significant poor survival for both BCSS and OS in patients with TNM stage II–III. Furthermore, in the subgroup of HER2⁻/HR⁺, HER2⁺/HR⁺, and triple-negative cancer, TCNP also gave noticeable prognostic value for predicting poorer BCSS and OS. Therefore, these results provided adequate evidence of decreased survival in patients with TCNP vs TBPQ, particularly for the various kinds of breast cancer subgroups.

This study has some potential implications for clinical practice. In screening, the examination of the axilla for TCNP should be stressed, and MRI may be a more practical imag-

ing method for accurate diagnosis. Inevitably, there are also some limitations in the present study. First, SEER, as a large database, may present the risk of coding error, despite its having undergone quality monitoring and integrity evaluation. Second, the subjects with treatment of "no/unknown" radiation or chemotherapy did not necessarily fail to receive these treatments or there may be missing information. Therefore, the SEER did not help to compare the outcomes by treatment received. Furthermore, important variables such as disease-free survival information are missing from the SEER database. Therefore, studies are needed to further clarify our findings.

Conclusion

We summarized the clinical characteristics of TCNP and demonstrated that the tumor site of central and nipple position was an independent prognostic factor and had potential implications for clinical practice. Patients with TCNP represent a cohort in whom studies of more focused staging procedures are required to assess LN metastases more carefully. Moreover, treatment programs should be studied to determine whether this disease presentation should receive a specifically tailored therapy.

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Author contributions

FJ, W-KX, and C-QY were the main authors of the manuscript. They were involved in the conception, design, and coordination of the study as well as in data analysis, interpretation of results, and drafting the manuscript. KW was in charge of all experimental procedures. MY, L-LZ, H-FG, Y-FL, TZ, M-YC, W-PL, W-JP, and X-SZ participated in the experimental procedures and revised critically the content of the manuscript. All authors contributed to data analysis, drafting and revising the article, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

Disclosure

The authors report no conflicts of interest in this work.

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