Retrospective Study

Macrometastasis at selective lymph node biopsy: A practical going-for-the-one clinical scoring system to personalize decision making

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Abstract

BACKGROUND
Axillary sentinel lymph node biopsy (SLNB) is standard treatment for patients with clinically and pathological negative lymph nodes. However, the role of completion axillary lymph node dissection (cALND) following positive sentinel lymph node biopsy (SLNB) is debated.

AIM
To identify a subgroup of women with high axillary tumor burden undergoing SLNB in whom cALND can be safely omitted in order to reduce the risk of long-term complications and create a Preoperative Clinical Risk Index (PCRI) that helps us in our clinical practice to optimize the selection of these patients.

METHODS
Patients with positive SLNB who underwent a cALND were included in this study. Univariate and multivariate analysis of prognostic and predictive factors were used to create a PCRI for safely omitting cALND.

RESULTS
From May 2007 to April 2014, we performed 1140 SLN biopsies, of which 125 were positive for tumor and justified to practice a posterior cALND. Pathologic findings at SLNB were micrometastases (mic) in 29 cases (23.4%) and macrometastasis (MAC) in 95 cases (76.6%). On univariate analysis of the 95 patients with MAC, statistically significant factors included: age, grade, phenotype, histology,
INTRODUCTION

The presence of axillary nodal involvement continues to be a major breast cancer prognostic factor. Complete axillary lymph node dissection (cALND) has a dual role both to ensure local control of the disease and to add staging information to adapt adjuvant treatment. The extent of cALND has evolved over time. In 1990, consensus from National Institute of Health concluded that the local treatment of axillary disease in operable breast cancer included at least the removal of axillary levels I and II[1]. In many European countries, cALND included all three Berg levels[2].

Axillary recurrence after cALND is infrequent[3-6], although treatment-related morbidity could be relevant both in the short- and long-term, including lymphedema, which occurs in 23 to 25% of women[7-11], pain, and decreased mobility of the arm[10, 12]. During the 1980s, the possibility to identify a regional lymph node that was the first lymph node drainage station for breast cancer began to be considered[13]. Years later, different studies established the utility of selective sentinel lymph node biopsy (SLNB) as an alternative to axillary dissection in patients with clinically negative lymph nodes (cN0)[14-16], with the advantage of decreasing the risk of lymphedema associated with cALND[9,10,17].

The presence of sentinel node involvement was associated with the need to complete the study and treatment of the axilla with cALND. In recent years, emphasis has been placed on the possibility of reducing the need of cALND in patients with positive SLNB without worsening the prognosis. The results of the ACOSOG Z0011[18] and IBCSG 23-01[19] trials suggest that cALND could be avoided when sentinel lymphovascular invasion, lymph-node tumor size, and number of positive SLN. On multivariate analysis, only lymph-node tumor size (≤ 20 mm) and number of positive SLN (> 1) retained significance. A numerical tool was created giving each of the parameters a value to predict preoperatively which patients would not benefit from cALND. Patients with a PCRI ≤ 15 has low probability (< 10%) of having additional lymph node involvement, a PCRI between 15-17.6 has a probability of 43%, and the probability increases to 69% in patients with a PCRI > 17.6.

CONCLUSION

The PCRI seems to be a useful tool to prospectively estimate the risk of nodal involvement after positive SLN and to identify those patients who could omit cALND. Further prospective studies are necessary to validate PCRI clinical generalization.

Key Words: Sentinel lymph node biopsy; Complete axillary lymph node dissection; Preoperative clinical risk index; Macrometastasis

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Core Tip: The role of completion axillary lymph node dissection (cALND) following a positive sentinel lymph node biopsy (SLNB) is being actively debated. Patients with a positive SLNB performed at our institution who also underwent a cALND were analyzed. Univariate and multivariate analysis of prognostic and predictive factors were used to create a Preoperative Clinical Risk Index (PCRI). The PCRI could help estimate the risk in removing extra positive nodes beyond the SLN in order to identify which patients could safely avoid cALND. Further prospective studies are necessary to validate clinical generalization for suggested tool.
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Aforementioned studies show wide variability regarding tumor burden in the sentinel node, including patients with isolated tumor cells (ITC), micrometastasis and macrometastasis making it impossible to definitively establish the effect of omitting cALND in patients with a higher tumor burden in the SLNB.

Our objective was to identify a subgroup of women with high axillary tumor burden in SLNB in whom cALND can be safely omitted in order to reduce the risk of long-term complications. The second endpoint is to create a Preoperative Clinical Risk Index (PCRI) that helps us in our clinical practice to optimize the selection of these patients.

MATERIALS AND METHODS

We retrospectively reviewed the experience of our Breast Cancer Unit with SLNB and cALND procedures from 2007 to 2014. This study received ethical approval from the Inhouse Local Ethics and Clinical Committee (Code: 16.04.0940-GHM) the April 27, 2016.

Inclusion criteria

All patients included in this analysis had breast cancer diagnosed by core needle biopsy. Axillary presurgical ultrasound with fine needle biopsy was done if required. Magnetic resonance imaging (MRI) of the breast was performed in all patients where it was technically feasible. Before 2012, SLNB was only done in patients with tumors less than 3 cm. However, since then, tumors smaller than 5 cm were also considered for SLNB. The sentinel lymph node assessment was performed intraoperatively, with hematoxylin-eosin and immunohistochemistry analyses deferred. Data from all patients with positive SLNB who underwent cALND were collected. From 2012, cALND was not carried out in the presence of micrometastasis.

Database information included the patients’ age, menopausal status, and laterality, as well as tumor characteristics at diagnosis (tumor size, number of foci, IHC, Her-2 status, grade, presence of intraductal carcinoma), surgical treatment data (surgery type, date, margins status, postoperative tumor size), and data of lymph nodes analysis (SLN, nodes from the accessory axillary fat and nodes from axillary dissection). In each of the sections, isolated tumor cells (ITC), micrometastasis (mic) and macrometastasis (MAC) were differentiated.

All patients with pathologic involvement of the axilla received adjuvant whole breast/chest wall radiotherapy together with comprehensive regional nodal irradiation (RNI) according to published international guidelines[21,22].

Statistical analysis

Patients’ information was obtained from the patients’ records, and collected data were subsequently analyzed with the SPSS software, version 17.0 (Released 2008. IBM SPSS Statistics for Windows. Version 17.0 Armonk, NY: IBM Corp).

Once the data had been validated and refined, the statistical analysis was carried out following this way: (1) Exploratory analysis of all selected variables; (2) Descriptive analysis of the sample: distribution of all variables; (3) Description of the clinical data: patients, tumor and type of surgery; (4) Description of sentinel node biopsy procedures (detailed pathological results of SLNs, axillary fat and axillary dissections); (5) Descriptive statistics of the patients with macrometastasis in the sentinel node; (6) Univariate analysis with the Chi-square test was performed to compare every clinical parameter with the variable “presence” or “absence” of positive nodes in the lymphadenectomy”. The parameters considered in the univariate analysis of the 95 cases of MAC were: age, menopausal status, laterality, tumor size, uni- or multicentricity, grade, phenotype, histology, association to in situ carcinoma, type of surgery, number of removed nodes, number of MAC, and the sum of MAC and mic; (7) For those significant variables in the univariate analysis, the Odds Ratio were calculated by using a logistic regression model using the Stepwise method to know which of them had the greatest impact on final results. The Pearson correlation coefficient was used. The level of significance used was α = 0.05 (95% confidence interval), therefore those differences whose P value < 0.05 were considered significant;
and (8) Finally, a numeric value was assigned to selected variables by calculating the logarithms of the results to develop a numerical tool, the PCRI. Different cut-off points of this tool, according to the cut-off point recommended by Krag et al[14] in the NSABP 32 trial, were simulated in order to find the one that left the least number of false negatives (low risk, FN must be < 10%, specificity of 90%).

RESULTS

From May 2007 to April 2014 a total of 1140 women underwent an SLNB procedure. One-hundred and twenty-five (11%) were positive for tumor and, hence, underwent posterior cALND. The average number of sentinel nodes dissected was 2.12 (1-5). Pathologic findings after SLNB were micrometastasis (mic) in 29 cases (23.4%) and macrometastasis (MAC) in 95 cases (76.6%).

We focused our analysis on patients with MAC. In our patients, the probability of harboring additional lymph node disease in the cALND in patients with MAC in SLNB was 36%.

Patients’ characteristic of the 95 patients with SLN macrometastasis are summarized in Table 1.

Univariate analysis

For the 95 patients with MAC, the parameters that showed statistical significance in the univariate analysis were: Age (cut point 40 years-old), grade, phenotype, histology, lymphovascular invasion, tumoral size, and number of positive sentinel nodes ($P < 0.05$ for all comparisons) (Table 2).

Multivariate analysis

When analyzing all the parameters that had been significant in the univariate analysis, only the tumor size less or equal to 15 mm and the presence of 1 affected SLN remained significant. The variable of up to 2 sentinel lymph nodes affected by MAC or mic as suggested by ACOSOG Z0011 was also studied. In our population, the presence of one MAC is the variable with the greatest statistical significance for predicting disease in the axillary dissection (Table 3). Of the 95 patients collected with MAC, 79 cases had only 1 affected SLN. For this reason, this data has been used as a cut-off point for the population analyzed.

PCRI

To avoid bias in the results due to the simultaneous presence of mic, a subgroup of 79 patients with exclusively one sentinel node with MAC in SLNB was selected from our original population, which served as the basis for developing the PCRI tool. Although having such a group of cases with little probability of involvement of other nodes in the cALND, these patients form a heterogeneous group. Thus, in the 79 women with only 1 affected SLN, we still found an additional 30.4% of disease in cALND, which is not an acceptable result to justify stopping cALND in all patients.

Based on the results of the univariate logistic regression of those variables that are related to the presence of additional disease on cALND (size, age, histology, grade and phenotype) in the group of 79 cases with presence of only one macrometastatic SLN, a numerical tool was created by giving each of the parameters a numeric value in order to be able to predict preoperatively which patients would not benefit from a cALND (Table 4). To facilitate the usability of the prognostic index, we decided to include only those factors that could be known before breast surgery. Appropriate numeric value was calculated by the resulting rate of involvement when considering this value in isolation and then taking these results to base 10 (Figure 1).

Numeric values assigned to age, grade, size, histological type, and phenotype were included. The resulting value of the PCRI was related to the risk of having other lymph nodes involved in the axillary dissection. According to the PCRI, patients could be divided into two groups based upon a cut-off point of 15.

Patients in the low-risk group (index score $\leq 15$, 42 patients) were at risk of having other affected nodes less than 10% and in whom the cALND would have added to morbidity with minimal extra information. Only 5% changed from the pN1a state after axillary surgery. Ninety-five percent of patients had 1 or 2 positive nodes in total. Patients in the high risk group (index score $> 15$, 37 patients) had a 54% of risk of having nodal involvement in the cALND. The pN1a stage changed in 13.5% of patients after lymphadenectomy. Seventy-three percent of the patients had 1 or 2 positive nodes in total, and 13.5% had 3 affected lymph nodes. For these patients, axillary
| Characteristics                              | n = 95 | %     |
|---------------------------------------------|--------|-------|
| Age (range), yr                             | 51.9 (23-82) | -     |
| Average tumor size, mm                      | 17.4   | -     |
| Tumor size                                  |        |       |
| 0-10                                        | 14     | 14.7  |
| 11-15                                       | 25     | 26.3  |
| 16-20                                       | 26     | 27.4  |
| 21-30                                       | 18     | 18.9  |
| 31-40                                       | 6      | 6.3   |
| > 50                                        | 1      | 1.1   |
| In situ                                     | 5      | 5.3   |
| Menopausal status                           |        |       |
| Pre                                         | 50     | 52.6  |
| Post                                        | 45     | 47.4  |
| Histology type                              |        |       |
| Ductal                                      | 76     | 80    |
| Lobular                                     | 15     | 15.8  |
| Others                                      | 4      | 4.2   |
| Phenotype                                   |        |       |
| Luminal                                     | 83     | 87.4  |
| Basal                                       | 9      | 9.5   |
| Her-2                                       | 3      | 3.2   |
| Grade                                       |        |       |
| I                                           | 16     | 16.8  |
| II                                          | 54     | 56.8  |
| I + II                                      | 70     | 73.7  |
| III                                         | 24     | 25.3  |
| Unknown                                     | 1      | 1.1   |
| Surgery                                     |        |       |
| Conservative                                | 63     | 66.3  |
| Mastectomy                                  | 32     | 33.7  |
| Number of SN removed                        |        |       |
| 1                                           | 35     | 36.8  |
| 2                                           | 28     | 29.5  |
| 3                                           | 20     | 21.1  |
| ≥ 4                                         | 12     | 12.6  |
| Nº of positive SN (Mic + MAC)               |        |       |
| 0                                           | 4      | 4.2   |
| 1                                           | 65     | 68.4  |
| 2                                           | 20     | 21.1  |
| 3                                           | 4      | 4.2   |
| ≥ 4                                         | 2      | 2.1   |
**DISCUSSION**

The American Society for Medical Oncology (ASCO) and National Comprehensive Cancer Network (NCCN) initially proposed in their clinical guidelines to always perform cALND in patients with positive SLNs\[23,24\]. However, studies had shown that 53% of patients with positive SLN did not have disease in the non-sentinel nodes after cALND\[25\], and that positive SLN was the only metastasis in 40% to 60% of cases\[14,26-29\]. In about half of the women undergoing cALND, surgery would not provide more prognostic information, although it would add a greater risk of complications derived from the procedure. Therefore, the current recommendations of international Guidelines and Consensus Statements on the locoregional treatment of breast cancer state the possibility of omitting cALND in the case of a positive SLNB is considered low risk, especially in those patients who are going to receive radiotherapy after surgery\[24,30\]. Omission of cALND is more frequently offered to patients who present lymph node involvement in the form of micrometastasis exclusively than in those patients with macrometastasis in the SLN\[30-32\].

When this study was conceived and designed, 2 large randomized trials had demonstrated the efficacy and safety of omitting cALND in select groups of patients\[7,33\]. Compared with what was observed by ACOSOG Z0011\[18\] and AMAROS\[7\] trials, our study presents very similar characteristics, both in the mean age and in the rate of ductal or lobular tumors, although in our series we found 8% more grade I/II tumors and half the cases of lymphovascular invasion (LVI) than reported in ACOSOG Z0011, which is considered a marker of aggressiveness and worse prognosis\[34\]. Finally, it should be noted that the tumor phenotype is known in all of our patients including hormone receptors, overexpression of the HER2 protein and the proliferation factor Ki-67. These data were not collected in the AMAROS; however, the ACOSOG Z0011 study collected data on hormone receptors, but not on Her2 or Ki 67, making a direct comparison difficult. With respect to the surgery performed, in ACOSOG Z0011, all patients received conservative surgery. In AMAROS, the number of conservative surgeries drop to 82%. In our study, 70% of the patients had conservative surgery while 30% received mastectomies. This could allow us to apply the
### Table 2 Univariate analysis of 95 patients with presence of sentinel lymph nodes with macrometastases

|                         | No additional metastatic lymph nodes | Additional metastatic lymph nodes | P value |
|-------------------------|--------------------------------------|----------------------------------|---------|
| Age                     |                                      |                                  |         |
| < 40                    | 66%                                  | 44%                              | 0.04    |
| ≥ 40                    | 32.6%                                | 67.4%                            |         |
| Hormonal status         |                                      |                                  |         |
| Premenopausal           | 70%                                  | 30%                              | NS      |
| Postmenopausal          | 60%                                  | 40%                              |         |
| T stage                 |                                      |                                  |         |
| T1                      | 75.8%                                | 24.2%                            | < 0.05  |
| T2                      | 47%                                  | 53%                              |         |
| T3                      | 0                                    | 100%                             |         |
| Tumor size              |                                      |                                  |         |
| ≤ 15 mm                 | 85%                                  | 15%                              | < 0.05  |
| > 15 mm                 | 49%                                  | 51%                              |         |
| Histologic type         |                                      |                                  |         |
| Ductal carcinoma        | 67.1%                                | 32.9%                            | NS      |
| Lobular carcinoma       | 47%                                  | 53%                              |         |
| Grade                   |                                      |                                  |         |
| I + II                  | 68.6%                                | 31.4%                            | NS      |
| III                     | 54%                                  | 46%                              |         |
| Molecular subtype       |                                      |                                  |         |
| Luminal A/B             | 65.6%                                | 34.4%                            | < 0.05  |
| Basal                   | 22.2%                                | 77.8%                            |         |
| HER2                    | 100%                                 | 0                                |         |
| LVI                     |                                      |                                  |         |
| Present                 | 66.7%                                | 33.3%                            | NS      |
| Absent                  | 61.5%                                | 38.5%                            |         |
| Number of positive SLN  |                                      |                                  |         |
| 1                       | 69.6%                                | 31.4%                            | < 0.05  |
| ≥ 2                     | 43.7%                                | 66.3%                            |         |
| ACOSOG Z0011 criteria   |                                      |                                  |         |
| ≤ 2 SLN+                | 68.5%                                | 31.5%                            | < 0.05  |
| > 2 SLN+                | 16%                                  | 84%                              |         |

LVI: Lymphovascular invasion; NS: Not significant; SLN: Sentinel lymph nodes.

The results of the SLNB and cALND are very similar in our study to those reported by AMAROS and ACOSOG Z0011. In our study, 69% of patients did not have additional metastatic involvement in cALND, similar to 72% and 73% of AMAROS and ACOSOG Z0011, respectively. Our study seeks to identify which patients will not have more axillary disease, even though the SLNB is positive for macrometastasis. The presence of macrometastasis in SLNB has been related to a higher probability of finding additional metastatic lymph nodes after cALND. According to other groups’ experience, the incidence of involvement in the cALND beyond the SLN was from 40 to 58% [23] when in the presence of MAC, while for mic it fell to 20% [23], and for ITC (isolated tumor cells), it
Table 3 Multivariate analysis of 95 patients with sentinel lymph nodes with macrometastases

|                          | P value | Odds ratio |
|--------------------------|---------|------------|
| Tumor size               |         |            |
| ≤ 15 mm vs > 15 mm       | 0.002   | 2.004      |
| Number of metastatic SLN |         |            |
| 1 vs > 1                 | 0.021   | 2.573      |
| ACOSOG Z0011 criteria    |         |            |
| ≤ 2 SLN+ vs > 2 SLN+     | 0.045   | 10.195     |

SLN: Sentinel lymph nodes.

Table 4 Univariate analysis of variables related to the presence of additional disease on completion axillary lymph node dissection in the group of 79 cases with presence of only one macrometastatic sentinel lymph nodes

| Additional metastatic lymph nodes, n = 79 |
|-----------------------------------------|
| Age                                     |
| < 40                                    | 50%   |
| ≥ 40                                    | 28%   |
| ≤ 15 mm                                 | 85%   |
| > 15 mm                                 | 49%   |
| Histologic type                         |
| Ductal carcinoma                        | 28%   |
| Lobular carcinoma                       | 50%   |
| Mucinous                                | 0     |
| Tumor size                              |
| ≤ 10 mm                                 | 7%    |
| 11-15 mm                                | 10%   |
| 16-20 mm                                | 42%   |
| > 20 mm                                 | 47%   |
| Grade                                   |
| DCIS                                    | 0     |
| I                                       | 18%   |
| II                                      | 31%   |
| III                                     | 45%   |
| Molecular subtype                       |
| Luminal A/B                              | 28%   |
| Basal                                   | 55%   |
| HER2                                    | 0     |

was roughly 12% (23). In our series, the probability of harboring additional lymph node disease in the cALND in patients with MAC in SLNB was 36%.

Currently, there are 3 ongoing trials to provide evidence in this regard. The SINODAR ONE trial[35] randomized patients with 1 or 2 macrometastasis in SLNB and treated with conservative surgery or mastectomy to receive either cALND remain under observation. Both groups received adjuvant therapy. The INSEMA trial[36] included patients with tumors T1-2 and conservative surgery or mastectomy and were randomized to undergo either SLNB remain under observation. Patients with a positive SLNB with 1 or 2 affected nodes are then re-randomized to exclusive SLNB or cALND. Finally, in the POSNOC trial[37], T1-2 breast carcinomas with 1 or 2 macrometastasis in SLNB are randomized after conservative surgery or mastectomy to either undergo axillary treatment with cALND or RNI or remain under observation. In both arms the indicated systemic treatment is added. The results of these 3 studies
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![Image](https://www.wjgnet.com)

Figure 1 Numeric values assigned to each variable to create an individualized Preoperative Clinical Risk Index.

| Risk index (PCRI) | Preoperative Clinical Risk Index (PRCI) ≤ 15/> 15 |
|-------------------|--------------------------------------------------|
| **Age**           | 2.8                                              |
| ≥ 40              | < 40                                             |
| **Histological type** |                                                  |
| Ductal            | Lobular                                         |
| 2.8               | 5                                                |
| **Phenotype**     |                                                  |
| Her2              | Luminal                                         |
| 0                 | 2.8                                             |
| Basal             | 5                                                |
| **Grade**         |                                                  |
| In situ           | GI                                              |
| 0                 | 2                                               |
| GII                | 3                                               |
| GIII               | 5                                               |
| **Tumor size (mm)** |                                              |
| ≤ 10              | 11-15                                           |
| 15                 | 16-20                                           |
| > 20               | 5                                                |

Figure 1 Numeric values assigned to each variable to create an individualized Preoperative Clinical Risk Index.

might elucidate the feasibility and safety of omitting cALND in patients with macrometastasis in SLNB.

Attempts have been made to develop tools to estimate the risk of finding additional positive lymph nodes beyond the SLN based on the presence of different factors: multifocality, presence of LVI, or the hormone receptors status. To date, there are currently seven published nomograms in use: the Memorial Sloan Kettering Cancer Center nomogram[38], the Cambridge nomogram[39], the Turkish Federation nomogram[40], the Stanford nomogram[41], the MD Anderson Cancer Center nomogram[42], the Tenon nomogram[43], and the MOU nomogram[44]. However, these nomograms do not seem to have good reproducibility in populations other than their own[45]. For a nomogram to be successful and clinically applicable, it must identify the majority of patients who might have non-sentinel node involvement, and the false negative rate should be ≤ 10%[45]. We have analyzed prognostic factors in our treated patients in order to design a personalized tool that, preoperatively and based on clinical and tumoral data, would help us to select patients with macrometastasis determined by SLNB with a risk of less than 10% of additional positive nodes in cALND, which would potentially allow them to omit this procedure regardless of the type of surgery and adjuvant treatments, similar to the initial approach of Sentinel Lymph Node Biopsy in NSABP. Our clinical index differs from other widely distributed nomograms, such as the Memorial Sloan Kettering Cancer Center nomogram (MSKCCn) and the MD Anderson Cancer Center nomogram (MDACCn) designed to infer involvement beyond the SLNB. One factor that is different in our clinical index is that our index takes into account the age of the patient, which is not included in either the MSKCCn or the MDACCn. However, tumor size is assessed identically in both the MSKCCn and the MDACCn, as well as in our PCRI. Histology and grade are assessed jointly in the MSKCCn, while MDACCn does not assess tumor grade. Both nomograms assess the total number of nodes removed: MSKCCn as negative sentinel nodes, and MDACCn as total nodes removed. The presence of multifocality was associated with a higher risk in the MSKCCn, but not in our index, and it was not assessed in the MDACCn. LVI is recorded in both nomograms, since it implies a worse prognosis. Our data support this result, but as it is not a parameter that we can obtain preoperatively, and therefore was not included in our index. Likewise, the extranodal extension included in the MDACCn is not a parameter that we can have preoperatively, so it has also not been included in our index. Finally, the tumor phenotype is not collected in the MDACCn and is only partially collected in the MSKCCn, which assesses the absence or presence of estrogen receptor in the tumor.

The most important strength is that our PCRI is that the information can be obtained preoperatively, allowing a personalized decision-making process that involves the patient before surgery. The biggest weakness of our study is the retrospective nature of the study, which limits the findings of the study. The population we assessed does not correspond to a specific geographic area, nor has participated in a standard screening program, making the findings less biased for a specific population. Our
patients come both from our own gynecology consultations, where opportunistic screening is performed following the American guidelines[23], as well as from other centers. However, the patients have private health insurance policies that could cause selection bias from the population. Additionally, comparing the characteristics of our population with the multicenter studies, American Z0011 and European AMAROS, we did not find any significant differences between the three patient populations. In addition, the phenotypes of basal and Her-2 were not as represented as other phenotypes in the total of patients, suggesting those patients should be specifically discussed in the Tumor Board.

CONCLUSION

More than half of our patients with the presence of MAC in the SLNB had no involvement of the non-sentinel nodes, and only 1 in 8 changed from the pN1a stage, which means that we have performed too many cALND on our patients.

The PCRI we outlined is based on a detailed analysis of the group of patients with MAC in the SLN, avoiding the biases of mic common in other trials, and is a useful tool to prospectively estimate the risk of nodal involvement beyond the positive SLN and to identify those patients who would benefit from a cALND afterwards.

Further studies are necessary to validate feasibility and accuracy of this PCRI.

ARTICLE HIGHLIGHTS

Research background
The role of completion axillary lymph node dissection (cALND) following positive sentinel lymph node biopsy (SLNB) is being actively debated.

Research motivation
Patients with positive SLNB performed at our institution who underwent a cALND were analyzed.

Research objectives
This study aims to create a Preoperative Clinical Risk Index (PCRI).

Research methods
Univariate and multivariate analysis of prognostic and predictive factors were used to create a PCRI.

Research results
PCRI could help estimate the risk of having extra positive nodes beyond the SLN in order to identify in which patients cALND could be safely omitted.

Research conclusions
Further prospective studies are necessary to validate clinical generalization for the suggested PCRI.

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