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Staging of the Axilla in Breast Cancer and the Evolving Role of Axillary Ultrasound

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Abstract: Axillary lymph nodes have long been recognized as a route for breast cancer to spread systemically. As a result, staging of the axilla has always played a central role in the treatment of breast cancer. Anatomic staging was believed to be important for two reasons: 1) it predicts prognosis and guides medical therapy, and 2) it is a potential therapy for removal of disease in the axilla. This paradigm has now been called into question. Prognostic information is driven increasingly by tumor biology, and trials such as the ACOSOG Z0011 demonstrates removal of axillary disease is not therapeutic. Staging of the axilla has undergone a dramatic de-escalation; however, sentinel lymph node biopsy (SLNB) is still an invasive surgery and represents a large economic burden on the healthcare system. In this review, we outline the changing paradigms of axillary staging in breast cancer from emphasis on anatomic staging to tumor biology, and the evolving role of axillary ultrasound, bringing patients less invasive and more personalized therapy.

Keywords: breast cancer, axillary staging, SLNB, ALND, axillary ultrasound, Z0011

Introduction to Staging of the Axilla in Breast Cancer

Breast cancer is the most prevalent cancer in women, and the second-leading cause of cancer death. In 2021, it is estimated that 280,000 cases of breast cancer will be diagnosed in the US, resulting in over 43,000 deaths.1 Axillary surgery has long been central to the staging and treatment of breast cancer. In the 19th century, Rudolph Virchow postulated that axillary lymph node metastases facilitated the distant metastatic spread of breast cancer.2 This led William Halsted to propose radical mastectomy (excision of the breast tissue, pectoralis muscle, and ipsilateral axillary lymph nodes) for the treatment of breast cancer.3 In the century since Halsted pioneered the radical mastectomy, advances in diagnostic imaging, systemic therapy, radiation therapy and an improved understanding of breast cancer biology have led to a dramatic de-escalation of breast cancer surgery.4 The modified radical mastectomy replaced the radical mastectomy. Breast conserving surgery further decreased the invasiveness of breast cancer surgery. Sentinel lymph node mapping made possible sentinel lymph node biopsy (SLNB) for early-stage breast cancers, sparing patients the morbidity of axillary lymph node dissection (ALND).5,6 Recent advances in axillary imaging and molecular profiling of primary breast cancers have called into question the necessity of SLNB/axillary surgery. Axillary ultrasound (AUS) has been proposed as a non-invasive and cost-effective method of staging the axilla.
alternative to SLNB for staging the axilla in breast cancer. Here, we review evolving paradigms for staging of the axilla in breast cancer, with a particular emphasis on the current and future role of axillary ultrasound.

Implications for Medical Decision Making

Axillary surgery has two potential roles in breast cancer management. First, it provides anatomic staging information which is used to predict prognosis and inform medical decision making related to adjuvant therapy. Second, it is a potential therapy for management of disease in the axilla. Both of these roles have been called into question, leading to an ongoing reevaluation of the necessity of axillary surgery.

The presence of axillary lymph node metastases has historically been considered an important prognostic factor in breast cancer, and impacted many aspects of medical decision making. Anatomic staging of breast cancer includes both clinical and pathological staging. Clinical staging integrates physical examination and imaging findings, while pathological staging incorporates the results of surgical pathology. In the AJCC TNM staging schema, metastatic disease in 1–3, 4–9, and ≥10 axillary lymph nodes constitutes N1, N2, and N3 diseases, respectively. Greater nodal involvement is associated with a worse prognosis, suggesting that more extensive surgery, radiation, and systemic therapy may be beneficial.

With improved understanding of tumor biology, medical decision-making is increasingly dependent on molecular profiling of the primary tumor. Biomarker and gene expression profiles are now routinely used to predict prognosis and response to therapy. Biomarker profiles are a surrogate for molecular subtype, and increasingly drive neoadjuvant and adjuvant systemic therapy decisions. Biomarker profile is particularly important for decisions regarding the use of targeted therapies, such as endocrine therapy, or therapy with HER2-targeting agents. Molecular profiling with tests, such as Oncotype Dx, provides additional prognostic information and is often more informative than anatomic staging information for predicting response to therapy.

Recent basic science studies using next-generation sequencing technologies confirm that most breast cancer metastases arise from the primary tumor rather than lymph node metastases. This is consistent with the hypothesis that the process of acquiring the biologic ability to metastasize (as outlined in the classic hallmarks of cancer) typically takes place at the primary tumor, not at lymph node metastases.

Therapeutic Considerations

As noted above, clinical decision-making paradigms have evolved in breast cancer and anatomic staging has decreased in importance in terms of medical decision-making. In addition, the therapeutic benefit of axillary surgery appears to be limited. The ACOSOG Z0011 trial randomized women with clinical T1-T2 N0 breast cancer and 1–2 positive sentinel lymph nodes at the time of SLNB to no further axillary surgery or completion ALND. Z0011 demonstrated no local control or survival advantage with completion ALND, suggesting that the therapeumatic benefit of axillary surgery is minimal or non-existent. Similar results were obtained in the IBSCG 23–01 trial. These trials have led to a paradigm shift in the perceived role of axillary surgery. Axillary surgery is now considered to be primarily a staging procedure, with minimal to no therapeutic benefit. This is reflected in the 16th St. Gallen International Consensus Guidelines in 2019 which states completion ALND is not indicated if patient receives post-lumpectomy radiation and appropriate systemic adjuvant therapy in early-stage breast cancer.

Current Modalities for Axillary Staging

Physical Examination

Axillary staging in newly diagnosed breast cancer begins with physical examination. However, physical examination has consistently been shown to lack sensitivity and specificity. Physical examination lacks the sensitivity for detection of microscopic or even subcentimeter axillary lymph node metastases. Specificity is complicated by the inability to distinguish reactive lymph nodes from lymph node metastases. Various studies have found that the positive predictive value ranges from 61 to 84%, and negative predictive value from 50 to 62%. Additional testing is required to accurately stage the axilla.

Axillary Ultrasound

Axillary ultrasound (AUS) is the most commonly used imaging modality for evaluation of the axilla in breast cancer. The evaluation is typically performed using a high frequency (7.5–17 MHz) linear-array transducer with the patient in supine oblique position, arm abducted.
and externally rotated with hand above head (“bathing beauty” position).23 AUS is non-invasive, affordable, and leverages readily available technologies. AUS has a PPV of 56–90% and NPV of 76–84%.22,24 Like all ultrasound technologies, however, it is limited by operator dependency and patient body habitus.

Sentinel Lymph Node Biopsy

Historically, axillary lymph node dissection (ALND) was performed routinely, providing staging information. In the 1990s, several randomized controlled trials demonstrated no difference in locoregional control, disease free survival, and overall survival between SLNB and ALND in patients with a negative SLNB.25,26 Furthermore, SLNB was associated with better quality of life, upper extremity function, and fewer side effects, such as chronic pain, lymphedema, and sensory deficits.5,6,27,28 In early-stage breast cancer, SLNB has become the standard of care for staging of the axilla. A meta-analysis of 69 trials with over 8000 patients found SLNB accurately mapped the sentinel node in 96% of patients with an average false negative rate of 7.3%.29

Axillary Lymph Node Dissection

ALND is now typically reserved for patients with significant axillary disease. It is primarily considered a therapeutic strategy in this context. In the aftermath of ACOSOG Z0011 and IBCSG 23–01 trial, ALND is still indicated for patients with three or more metastatic lymph nodes.30 ALND is also indicated in those with less than three positive nodes who either refuse or are not candidates for adjuvant radiation. Since the Z0011 trial excluded patients with extranodal extension, ALND is still performed for patients with less than 3 positive SLNs but with >2 mm extranodal extension up to 33% may have more extensive disease involving more than 4 nodes.31 As mentioned previously, ALND is associated with significant morbidity and thus patient selection must be carefully considered.

Biologic Staging

Recent decades have seen a dramatic acceleration in the understanding of breast cancer biology and the concomitant development of targeted therapeutics. Biomarker profile and gene expression profiling assays, such as OncotypeDx, are currently used to drive clinical decision making. The increasing role of tumor biology is reflected in the 8th edition of the AJCC staging system. The AJCC system now combines anatomic staging information with HER2 and hormone receptor status to create a clinical prognostic stage. A pathologic prognostic stage was further created with the addition of histologic grade and OncotypeDx RS score.32 The revised AJCC staging system more accurately predicts prognosis by integrating both anatomic and biologic staging information. The revised AJCC staging system was derived from over 300,000 patients enrolled in the National Cancer Database between 2010 and 2012 representing over 70% of breast cancers diagnosed in the US.32 However, the prognostic validity of the revised AJCC system assumes availability and access to recommended adjuvant therapy. For example, HER2+ tumors are downstaged because there is an effective targeted therapy. Similarly, the incorporation of OncotypeDx RS score assumes patients will be managed appropriately.

Current Staging Algorithms

The current algorithm for staging of the axilla in early stage (T1-2) breast cancer starts with clinical evaluation based on physical exam and imaging (summarized in Figure 1). Clinically node-negative patients typically undergo SLNB. Patients with negative sentinel nodes or only micrometastases do not need further axillary surgery.

Clinically node-positive patients are candidates for ultrasound-guided FNA or core needle biopsy (CNB). In the event of a negative FNA/CNB, SLNB is indicated. If FNA/CNB is positive, ALND is indicated unless neoadjuvant therapy is performed. In patients with positive FNA/ CNB who subsequently receive neoadjuvant chemotherapy and become clinically node negative, SLNB may be considered.

Axillary Ultrasound

Characteristics of Abnormal Lymph Nodes on Ultrasound

AUS can identify disease in ALN based on the presence or absence of defined sonographic criteria, such as absent or infiltrated hilum, or focal or diffuse cortical thickening (Figure 2).33–36 Normal lymph nodes in ultrasound have uniform hypoechoic cortex, central fatty hilum, and smooth margins.37 Any deviations from this normal appearance are potential signs of metastasis. A cortical thickness of greater than 3 mm is associated with more than three-fold increase in metastatic disease, loss of fatty hilum is associated with a 27-fold increase,
and tumor margin irregularity is associated with a 3-fold increase.\textsuperscript{38,39} Lymph node diameter greater than 10 mm is also associated with increased risk of metastatic disease. While these features have high odds ratios, individually they have poor AUC in predicting the presence of metastatic disease.
Axillary Ultrasound at Initial Presentation
Since physical examination is neither sensitive nor specific in terms of predicting the likelihood of nodal metastasis, axillary ultrasound (AUS) is an important non-invasive adjunct to the assessment of the axilla on initial presentation. Currently, AUS is most informative for the evaluation of patients with clinically palpable nodes. This offers the opportunity to perform FNA or core biopsy on suspicious nodes, helping to identify candidates for ALND rather than SLNB to streamline care. The ability of US FNA to predict disease was shown to have a sensitivity and specificity of 79.6% and 98.3%, respectively, in a meta-analysis by Houssami et al.40

The role of AUS in the pre-operative evaluation of the axilla has been called into question since the ACOSOG Z0011 trial. Some have argued that a positive AUS would unnecessarily commit many patients to ALND who met criteria for observation in the Z0011 trial, namely T1-2 tumors with 1–2 positive SLNs. However, there is evidence that disease identified by AUS FNA suggests a higher axillary disease burden compared to the disease identified by SLNB. Positive AUS FNA is associated with more positive nodes, larger tumors, higher tumor grades, extranodal extension, lymphovascular invasion, and distant metastasis. Several case series have found that only 3–5% of patients with positive AUS FNA met Z0011 criteria and could have avoided ALND, thus streamlining axillary surgical care for 28.6% of node-positive patients. This suggests that AUS continues to play a vital role in the pre-operative evaluation in newly diagnosed breast cancer.

Axillary Ultrasound After Neoadjuvant Chemotherapy
Historically, patients with node-positive disease underwent ALND after neoadjuvant chemotherapy (NAC). However, this approach is morbid, and 40% and 70% of patients convert from node-positive to node-negative disease following NAC and targeted therapy, respectively. Several prospective studies have looked into the efficacy of SLNB post-NAC. In the NSABP B27, SENTINA, ACOSOG Z1071, and SN FNAC trials, SLNB was limited by high false negative rates of 10.7%, 14.2%, 12.6% and 8.4%, respectively. Patients with a suspicious AUS were node positive 71.8% of the time compared to 56.5% for patients with a normal AUS. This suggests a role for AUS evaluation in order to guide post-NAC surgical management of the axilla. In a retrospective comparison of pre- and post-NAC imaging, AUS had a sensitivity of 69.8% for detecting persistent LN metastasis, which outperformed MRI and PET-CT.46

Furthermore, the utility of axillary ultrasound may depend on the molecular subtype of the cancer. Di Micco et al found PPV to be 100% and 70.5% for luminal A and luminal B breast cancer, respectively, and NPV to be 93.3% for both HER2 and triple negative breast cancers. Thus, AUS could potentially decide the need for ALND when used as part of a clinical decision algorithm that incorporates tumor biology.

As outlined in section 2.6, all patients with clinically positive axilla should undergo ultrasound evaluation. This serves not only as an opportunity for biopsy but also as a baseline imaging comparison in case NAC is used. The use of AUS in restaging of the axilla post-NAC varies by institution. When used, patients with previously clinically positive axilla who have downstaged to cN0 after NAC undergo AUS. If AUS demonstrates no persistent disease, patient is eligible for SLNB instead of ALND (Figure 3).

Axillary Ultrasound vs Sentinel Lymph Node Biopsy
Limitations of Sentinel Lymph Node Biopsy
Although the sentinel lymph node hypothesis is elegant, SLNB has important limitations suggesting that alternative approaches to axillary staging should be considered. First, SLNB is an invasive procedure, but is not considered

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**Figure 3**: Restaging of the axilla after NAC. ALND has traditionally been recommended for patients with a clinically positive axilla and positive FNAC/NB. For patients who are treated with NAC, evaluation with AUS can help reduce the utilization of ALND and concurrent patient morbidity. Following NAC, AUS can be used to assess response to therapy. If the AUS is negative after NAC, SLNB is recommended with ALND reserved for patients with ≥2 positive sentinel lymph nodes.
therapeutic. Second, the importance of anatomic staging information is eroding in an era where biomarker profile and gene expression profile are driving medical oncology decision making. Finally, SLNB represents an enormous investment in health care resources.

SLNB is an invasive surgical procedure with significant risks. Large prospective trials, such as ACOSOG Z0010, NSABP B-32, and the ALMANAC trial, have documented SLNB complications including allergic reactions to isosulfan blue dye (0.1–1.0%), wound infection (1.0–10%), seroma (7.1%), paresthesias (8.6–11%), and hematoma (1.4%).26,27,48,49 For many breast cancer patients, SLNB is the most morbid component of breast conserving surgery. Of note, not all SLNBs are successful. In 3–5% of cases, the SLNB cannot be identified, in which case current guidelines recommend that an ALND be performed.30 ALND is a morbid procedure with no therapeutic benefit in patients with clinical N0 disease.

Furthermore, SLNB currently represents an enormous investment in healthcare resources. Henry-Tillman et al found charges associated with SLNB at University of Arkansas to total $23,956.50 Based on this number, we estimate that the health care system invests approximately $3 billion dollars annually on an invasive surgical procedure that is not considered therapeutic, and has limited impact on medical oncology and radiation oncology decision making. McEvoy et al have conducted cost-effective analyses in T1-2 N0, HR+/HER2- breast cancer patients, demonstrating observation was superior to SLNB with net monetary benefit of $655,659 for observation vs $641,778 for SLNB. This suggests that the benefit of observation is not only economic but also reflected in improved quality of life.51

**Evolving Paradigms: Biologic vs Anatomic Staging**

In the post-Z0011 era, the centrality of surgical axillary staging to the management of breast cancer is being questioned. First, axillary surgery is not considered therapeutic. The ACOSOG Z0011 trial randomized women with clinical T1-T2 N0 breast cancer and a positive SLNB to no further axillary surgery or to completion ALND, and demonstrated no local control or survival advantage with completion ALND. Similar results were obtained in the IBSCG 23–01 trial.15,16 Second, the importance of anatomic staging information provided by SLNB is decreasing in importance. Decisions regarding adjuvant therapy are complex, and integrate a significant amount of clinical and pathologic information, such as age, performance status, biomarker profile, and gene expression profile.52–55 Although anatomic staging is important, it is decreasing in importance as a driver of medical decision making and this reality is beginning to be reflected in clinical guidelines.

Systemic therapy is increasingly driven by tumor biology, such as gene expression and receptor status. For instance, HER2+ breast cancers are at least considered for systemic chemotherapy regardless of nodal status.8 For HR+/HER2- cancers, chemotherapy is determined by nodal status only if there are ≥4 positive nodes, with gene expression assays used to determine the need for chemotherapy for cancers with less than 4 positive nodes. Finally, in triple negative breast cancer, chemotherapy is considered regardless of nodal status.

Adjuvant locoregional therapy is still driven by anatomic staging. Guidelines recommend additional regional radiotherapy when there are axillary lymph node metastases.8 However, this is increasingly called into question for early-stage breast cancer.56–58 Whelan et al, in a trial where women with node-positive or high-risk node-negative breast cancers were randomized to whole-breast radiation vs whole-breast radiation plus regional nodal radiation, found reduced recurrence but no improvement in overall survival after 10 years with nodal radiation.57 This suggests that future locoregional therapy should not only take into account anatomic staging but also tumor biology.

**Current Trials Comparing AUS to SLNB**

The feasibility of AUS as an independent staging modality is currently being explored. In a prospective pilot study, Cyr et al demonstrated AUS to be a promising non-invasive alternative to SLNB in patients with early-stage breast cancer.59 Three large clinical trials are currently underway to investigate the role of AUS in staging of the axilla in patients with early-stage breast cancer undergoing breast conservation surgery (BCS) with a clinically negative axilla (Table 1). These studies have the potential to dramatically alter the paradigm for axillary staging as the unifying hypothesis for these studies is that most patients with a negative AUS do not require additional axillary staging.

**SOUND Trial**

The SOUND Trial (Sentinel node vs Observation after axillary UltrasouND) is a prospective randomized
| Trial             | Type                                | Arms                                      | Endpoint                                      | Inclusion                                                                                     | Exclusion                                                                 |
|------------------|-------------------------------------|-------------------------------------------|-----------------------------------------------|------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------|
| SOUND (NCT02167490)  
N=1560  
(780 per arm) | Multicenter prospective randomized non-inferiority trial  
(Δ=2.5%) | Arm 1: SLNB ± ALND  
Arm 2: Observation | 1°: DDFS  
2°: Distant recurrences, axillary recurrences, disease free survival, overall survival, and quality of life measures | • Breast cancer ≤2 cm  
• cN0 (negative AUS or US FNA)  
• Candidate for BCS | • Synchronous distant metastases  
• Previous malignancy  
• Bilateral breast cancer  
• Multicentric or multifocal disease  
• Previous primary systemic therapy  
• Pregnancy or breast feeding  
• Pre-operative pathological diagnosis of axillary node metastasis  
• Pre-operative radiological evidence of suspicious nodes  
• Psychiatric hx |
| INSEMA (NCT02466737)  
Goal of N=7095 | Multicenter prospective randomized non-inferiority trial | 1st random:  
Arm 1: Observation  
Arm 2: SLNB  
2nd random (1-3 positive SLN):  
Arm 1: Observation  
Arm 2: Completion ALND | Invasive disease free survival (IDFS) | • Breast cancer ≤5 cm  
• cN0 (negative AUS or US FNA)  
• M0  
• Planned BCS | • Hx of malignancy in past 5 years  
• Non-invasive breast cancer  
• T3/T4 tumors  
• Neoadjuvant chemotherapy  
• Pregnant or lactating  
• Multicentric tumors  
• Planned intra-op RT or post-op partial breast RT alone |
| BOOG 2013-08 (NCT02271828)  
Goal of N=1644 | Multicenter prospective randomized non-inferiority trial | Arm 1: SLNB ± ALND  
Arm 2: Observation | Regional recurrence rate (up to 10 years) | • Invasive breast cancer  
• T1-2 tumors  
• cN0 with negative AUS or US guided FNA  
• Planned BCS | • cN+ disease  
• Bilateral breast cancer, metastatic disease  
• History of invasive breast cancer  
• Previous axillary surgery or RT  
• Pregnant or nursing  
• Prior malignancies within past 5 years |

**Abbreviations:** SLNB, sentinel lymph node biopsy; ALND, axillary lymph node dissection; AUS, axillary ultrasound; US, ultrasound; FNA, Fine Needle Aspirate; BCS, breast conservation surgery; RT, radiation therapy.
multicenter non-inferiority trial designed by the European Institute of Oncology of Milan (NCT02167490). Inclusion criteria include breast cancer ≤ 2 cm, clinically negative axilla with negative AUS or US guided FNA, and candidate for BCS. The trial has recruited 1560 patients who were randomized to either SLNB ± ALND or observation. The primary endpoint is distant-disease-free survival. Secondary endpoints are incidence of distant recurrences, axillary recurrences, disease-free survival, overall survival, and quality of life measures.

**INSEMA Trial**

Another trial underway is the multicenter German/Austrian Intergroup-Sentinel-Mamma (INSEMA) trial (NCT02466737). It is a prospective randomized non-inferiority trial with the goal of enrolling 7095 patients. The trial has two randomizations. The first will be to have no axillary surgery or SLNB. Patients who have undergone SLNB with 1–3 micrometastases in sentinel lymph nodes will undergo a second randomization for either no further axillary surgery or completion ALND. Inclusion criteria are women with breast cancer ≤ 5 cm, cN0 with negative AUS or US FNA, no evidence of metastasis, and planned BCS. Patients with a history of malignancy within last 5 years, non-invasive breast cancer, T3/T4 tumors, NAC, pregnant or lactating, multicentric tumors, and planned intraoperative radiotherapy (eg Intrabcam) or postoperative partial breast irradiation (eg multicatheter technique) alone are excluded. The primary outcome is disease-free survival after breast conservation surgery.

**BOOG 2013-08**

Finally, the Dutch BOOG 2013-08 (NCT02271828) is another non-inferiority multicenter randomized control trial that will randomize patients to observation or SLNB ± ALND. Inclusion criteria include pathologically confirmed invasive breast cancer, T1-2 tumors, cN0 with negative AUS or US guided FNA, and planned BCS. Patients with clinically positive nodes, bilateral breast cancer, metastatic disease, history of invasive breast cancer, previous axillary surgery or RT, pregnant or nursing, and prior malignancies within past 5 years are excluded. Primary outcome is a regional recurrence rate (up to 10 years). Unlike the SOUND and INSEMA trials, this trial will include patients who have received neoadjuvant chemotherapy.

**Future Staging Algorithms**

The completion of the SOUND, INSEMA, and BOOG 2013-08 trials will likely alter staging algorithms for T1-2 breast cancer with axillary ultrasound playing a central role. Evaluation of newly diagnosed T1-2 breast cancer will first undergo AUS (Figure 4). Patients with no suspicious findings are spared from additional axillary staging procedures and are eligible for observation. Those with suspicious findings will undergo ultrasound-guided FNA or core biopsy. If ultrasound-guided biopsy is positive, patient will proceed with the standard surgical staging algorithm of SLNB with or without completion ALND. Patients with a negative biopsy may be safely observed.

**Figure 4** Possible future algorithm for staging of the axilla in early stage breast cancer. Several studies are currently ongoing evaluating the ability of AUS to accurately exclude disease in the axilla. If these studies are successful, the paradigm for axillary staging will evolve. Patients with newly diagnosed T1-2 breast cancer will be evaluated with an AUS. Patients with negative AUS will not require further axillary staging. Patients with a positive AUS will undergo FNA/CNB. Patients with a negative FNA/CNB can be observed. Patients with a positive FNA/CNB should undergo SLNB, possible ALND.
New Applications and Technologies
Contrast Enhanced Ultrasound

Contrast enhanced ultrasound (CEUS) is a relatively new technique for imaging of the axilla introduced by Omoto et al. The current technique involves an intradermal injection of 1 cc of 25% albumin solution as contrast in the upper outer quadrant of the breast adjacent to the nipple. The breast is then massaged to encourage uptake of microbubbles into the lymphatic system. Using contrast pulse sequencing, microbubbles are followed in real-time to the axilla. The first axillary lymph node to be filled by contrast is assumed to be the sentinel node.

Xie et al described three types of lymph node enhancement: type I, homogenously enhanced; type II, heterogeneous enhancement; type III, weakly enhancing or non-enhancing. When type I nodes were considered as negative and type II & III nodes as positive, the authors found ultrasound to have a sensitivity and specificity of 86.2% and 84.7%, respectively, for detecting metastatic lymph nodes.

Current methods for identification of the sentinel lymph node during surgery include intraoperative use of lymphazurin/methylene blue dye and radiolabeled colloid. CEUS has been investigated as an alternative in identifying the target node. Comparing CEUS to methylene blue, Xie et al found no difference in correct identification of the sentinel lymph node between the two techniques. In a prospective case series of patients who underwent pre-operative CEUS-guided core needle biopsies followed by surgical staging, CEUS correctly identified the sentinel node in 555 out of 605 cases.

Another technique uses a dual-modality photoacoustic and ultrasound imaging system to directly identify methylene blue accumulation in lymph nodes. Uribe et al demonstrated this concept in a pilot study of 16 patients. The goal of these technologies is to eventually perform percutaneous sentinel node biopsies using these technologies as detailed below.

Ultrasound-Guided Biopsy of the Sentinel Node

Accurate identification of the sentinel lymph node offers the opportunity for percutaneous biopsy, which avoids the morbidity of an SLNB. In a meta-analysis, ultrasound-guided FNA and CNB had sensitivity of 74% and 88%, respectively, for diagnosing nodal metastases, and both had specificity of 100%. Some have proposed that CEUS-guided SLN-FNA can effectively replace SLNB in identifying patients requiring ALND, but there may be higher rate of false-negative results in HER2-negative patients. When used as a pre-operative screening tool, Deurloo et al showed a 14% decrease in the number of SLNBs required. However, some evidence suggests that in patients meeting ACOSOG Z0011 criteria, using US-guided FNA as pre-operative triage may lead to overtreatment.

Prediction Nomograms

Nomograms that incorporate ultrasound findings with tumor characteristics have been developed to predict the presence of lymph node metastasis. Qiu et al created the Shantou Nomogram assigning points to sonographic factors, such as lymph node diameter, cortical thickness, and fatty hilum loss, and tumor-related factors, including tumor size, histologic grade, and estrogen receptor status. The AUC was 0.9375 and 0.864 in the modeling group and validation group, respectively, confirming that nomograms may improve the ability of AUS in patient selection for axillary staging procedures. Tran et al showed that SLNB may be selectively omitted in patients with low probability of nodal metastasis based on the Shantou nomogram if used as a pre-operative screen.

Intraoperatively, a nomogram could be used to predict the presence of lymph node metastasis using ex vivo shear wave elastography (SWE), an ultrasound modality. In a cohort 55 patients, excised lymph nodes were first examined using SWE and subsequently sent for pathological analysis. AUC was 0.856 and 0.791 in the development and validation cohorts, respectively.

Targeted Axillary Lymph Node Dissection

Post-NAC, SLNB has been shown to have lower yield and higher false-positive rates. Targeted axillary lymph node dissection (TAD) has been proposed, which involves the post-NAC removal of a node with documented metastasis on pre-NAC imaging. Ultrasound is used to mark diseased nodes pre-NAC with clips, ink, radiofrequency tags, or radioactive or magnetic seeds. The various techniques and their pros and cons are reviewed extensively by Banys-Paluchowski et al and clinical trials are currently underway to investigate their utility in clinical practice.

Radiomics

Radiomics is the process of converting radiologic images into high dimensional mineable data. After volume of
interest (VOI) is selected, extraction algorithms are used to generate datasets based on radiomic features (eg intensity, shape, texture, or wavelet). \textsuperscript{79} Several nomograms have been created using ultrasound radiomics and LASSO regression to predict the presence of lymph node metastasis with high AUCs. \textsuperscript{80,81} With the application of deep learning to ultrasound radiomics datasets, known as deep learning radiomics ultrasonography (DLRU), axillary lymph node status may be accurately predicted with AUC of >0.90. \textsuperscript{82,83} Although radiomics is still in its infancy, it has great potential to pave the way for the personalization of surgical management of the axilla and “virtual biopsies.”

\section*{Conclusion}
\subsection*{Paradigm Shift from Anatomic Staging to Tumor Biology}
Axillary surgery in breast cancer has undergone a dramatic change – from axillary lymph node dissection to the use of sentinel lymph node biopsy, reserving full dissection only for select patients. Historically, axillary surgery offered important anatomic staging information and was believed to be therapeutic by removing what was considered to be an important nidus of future metastasis. However, medical decision-making is increasingly driven by data derived from the primary tumor such as biomarker and molecular profile. \textsuperscript{53,54} Furthermore, new information about the biology of tumor metastasis suggests that metastatic disease typically originates in the primary tumor rather than in metastatic lymph nodes. \textsuperscript{12-14} The ACOSOG Z0011 trial provides strong evidence that axillary surgery is no longer therapeutic, demonstrating that patients with early-stage breast cancer and 1–2 positive sentinel lymph nodes did not benefit from further axillary lymph node dissection. Taken together, evolving paradigms in breast cancer management have called into question the necessity of surgical axillary staging, and have mirrored broader changes in oncology to more personalized therapy informed by biomarkers. \textsuperscript{84-86}

\section*{Axillary Ultrasound in New Staging Algorithm}
Axillary ultrasound is an attractive modality to continue the de-escalation of the management of the axilla in breast cancer. Surgical staging is invasive for patients, and represents a significant economic cost for the healthcare system. Staging with AUS in lieu of surgical staging has demonstrated increased QALYs and monetary savings. \textsuperscript{51} Three multicenter prospective randomized clinical trials, the SOUND, INSEMA, and BOOG 2013–08, are underway to investigate the potential of using axillary ultrasound to stage early breast cancer. These trials will provide valuable information in constructing future staging algorithms where AUS will play a central role, potentially replacing surgical staging. Patients will benefit from avoidance of an invasive surgery, while the healthcare system will be relieved of an economic burden.

\section*{Outlook}
As AUS play a more central role in staging of the axilla, new technologies are on the horizon with the potential to increase its utility and expand its role. Biopsies guided by contrast enhanced ultrasound represent another potential alternative to SLNB. Prediction nomograms are simple tools that synthesize clinical, biological, and sonographic features to predict axillary disease burden and aid surgical decision making. \textsuperscript{71,72} The budding field of radiomics applies neural networks and deep learning to ultrasound imaging to construct models not possible previously. \textsuperscript{87} These developments could allow AUS to be incorporated into the treatment of advanced stage cancers beyond those included in the Z0011 trial. The surgical management of breast cancer has undergone many transformations over the past century. Axillary ultrasound represents the next phase in this radical transformation, and is certain to play a central role in future staging algorithms of the axilla in breast cancer.

\section*{Author Contributions}
Michael Y Chen, MD, William E. Gillanders, MD
All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal; gave final approval for the version to be published; and agreed to be accountable for all aspects of the work.

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