Ultrasound (US) image-guided core biopsy of the breast

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Abstract: Results of partial mastectomy or lumpectomy including margin negativity are improved when preoperative diagnosis is obtained. This article describes the various techniques, instruments, utility and complications of the techniques. Emphasis is given to ultrasound (US) usefulness and effectiveness.

Keywords: Core needle biopsy (CNB); vacuum-assisted biopsy; excisional biopsy; breast cancer; breast surgery

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Recent trends in breast cancer treatment have favored breast conservation surgery with an emphasis on improved cosmesis. In the last twenty years of breast surgery, one of the most important advances has been the ability to diagnose breast cancer outside of the operating room utilizing the techniques of percutaneous core needle biopsy (CNB), stereotactic biopsy, and vacuum-assisted biopsy techniques. As nearly 80 percent of mammographic abnormalities are benign, the need for open biopsy has dramatically decreased, in turn supporting the use of minimally invasive percutaneous techniques. Utilizing excisional biopsy for diagnostic purposes often requires repeat surgery to establish clear margins in cases of a cancer diagnosis, propelling a shift towards percutaneous biopsy (1-5). The ability to obtain a diagnosis of cancer prior to surgery can allow for proper pre-operative planning, decrease the subsequent positive margin rate, and thus decrease the re-excision rate.

Percutaneous CNB is the preferred minimally invasive technique for diagnosing both palpable and non-palpable lesions. An international interdisciplinary consensus conference held in 2001, 2005, and again in 2009 agreed that percutaneous biopsy of breast lesions should be the gold standard biopsy method with use of US guidance if the lesion is amenable and stereotactic biopsy for calcifications not visualized on US (6). The Agency for Healthcare Research and Quality published an evidence report in 2014, that included 160 studies, and concluded that women were 15 times more likely to have their cancer treated with a single surgical procedure if they underwent image guided biopsy rather than open excisional biopsy (7). Breast surgeons are increasingly gaining expertise in breast US and US-guided core needle biopsies (CNB) (2,4). Surgeon-performed breast biopsies have shown to be cost-effective and associated with high patient satisfaction rates (3). Surgical societies are increasingly providing training programs and are certifying surgeon competency in these areas (2,4). In the United States, percutaneous CNB has nearly replaced fine needle aspiration (FNA) as the pre-operative diagnostic method of choice for breast lesions, as it provides a more definitive histological diagnosis and adequate tissue for prognostic markers (5-9). Performance of percutaneous biopsies for tissue diagnosis allows for optimization of surgical planning with concomitant staging of the axilla, therefore, decreasing the need for re-operation.
US-directed biopsy is performed for lesions that are palpable, but also for non-palpable, mammogram-detected lesions with a corresponding abnormality visualized on US. US-guided core biopsy is technically easier than stereotactic-guided biopsy as real-time imaging allows the surgeon to visualize the biopsy as it occurs. Using sterile technique and local anesthesia, a small puncture is made with an 11-knife blade and the needle is placed near the edge of the lesion and fired, which inserts it into the lesion for a sample to be retrieved. When placed parallel to the needle the position of the needle can be visualized on US (Figure 1).

A 12- or 14-gauge needle spring-loaded large core biopsy gun is used to remove several cores of tissue and in some instances completely remove the lesion by manually moving in and out of the insertion site, although automated core biopsy guns are also available. It is recommended that 5–10 cores be taken from a single lesion for adequate sampling.

If the core biopsy result is benign and is concordant with imaging findings, continued surveillance is acceptable. If the result is indeterminate or image-discordant, surgical excision is indicated to rule out malignancy. In order to determine the appropriate management and surveillance of a lesion, the histologic, imaging, and clinical findings must be taken into account for an assessment of concordance to be performed. When benign histology is obtained via CNB from a lesion that is concerning for malignancy either on imaging or on clinical exam then the biopsy is considered discordant. Discordance necessitates additional evaluation of such lesions by methods including repeating percutaneous biopsy or proceeding with surgical excisional biopsy (10). In addition, surgical excision is indicated for a core biopsy that demonstrates atypical hyperplasia (lobular or ductal) or lobular carcinoma in situ or neoplasia as the incidence of coexisting ductal carcinoma in situ or invasive carcinoma may be as high as 30% due to potential sampling error (11-14).

**Image-guided vacuum-assisted core biopsy**

Vacuum-assisted core biopsy is based on the same general principle as the CNB but represents a significant advance in technology (15-17). Vacuum is used to pull tissue into a sampling chamber, where it is removed with high-speed internal rotating knives. The specimen is then suctioned to a chamber outside the breast, where it can be retrieved. Multiple samples can be removed through this single-insertion technology, which has been approved for complete removal of benign imaged abnormalities under US or stereotactic guidance (Figure 2).

Image-guided vacuum-assisted biopsy (8- or 11-gauge) uses single-insertion technology with vacuum assistance for removal of imaged abnormalities via a suture-less incision (Figure 3).

Standard core needle (12- & 14-gauge) biopsy limits the amount of tissue available to pathologists for establishing the histologic diagnosis and biochemical markers. For example, a standard core biopsy specimen may show ductal

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**Figure 1** Core needle biopsy with ultrasound guidance. (A) Spring loaded, hand held core needle biopsy device; (B) insertion of the core needle into a lesion using ultrasound guidance to obtain core biopsy specimens of the tissue.
carcinoma in situ, but the final pathology of the lumpectomy specimen may demonstrate focal areas of infiltrating carcinoma. The vacuum-assisted core biopsy systems that are available provide for a more accurate diagnosis because larger specimens are removed than are possible with core biopsy and, in the case of image-guided vacuum-assisted excisional breast biopsy (IVEB), benign lesions are fully excised via removal of multiple cores. There are a plethora of vacuum-assisted devices available (Table 1).

Vacuum-assisted core biopsy offers the ability of obtaining larger (3–5 mm × 19 mm) contiguous samples from the same area by rotating the device rather than by withdrawal and reinsertion, as is necessary with the core biopsy needle. Theoretically, this minimizes seeding of the core tract and affords more accurate diagnosis (18,19). Patients diagnosed by vacuum-assisted or standard CNB have shown no difference in recurrence rates compared with patients diagnosed by excisional breast biopsy, suggesting that with radiation limited seeding of the needle tract does not affect outcome (20).

**Image-guided vacuum-assisted excisional breast biopsy (IVEB)**

The utility of the vacuum-assisted breast biopsy device under stereotactic guidance is well established (21,22). We hypothesized that the hand-held device could be used for diagnostic purposes as well as for therapeutic management (23). We facilitated the complete removal of benign-appearing lesions under US guidance with minimal complications and better cosmesis. Patients enrolled in
this study underwent an US-guided minimally invasive excisional breast biopsy through a 3-mm incision. Complete removal of the imaged abnormality was accomplished with a hand-held 8- or 11-gauge image-guided vacuum device (Mammotome®, Ethicon Endo-Surgery, Cincinnati, OH, USA), which uses a high-speed rotating cutter to remove intact cores of breast tissue (Figure 4). Cores measuring 3–5 mm × 19 mm were removed until the visualized abnormality was completely excised. The gauge of the device was determined by lesion size. Lesions 1.5 cm² and larger were excised with the 8-gauge device. If smaller lesions were removed, the 11-gauge mammotome device was used.

A total of 101 lesions were excised. The average size of the lesions was 1.9±1.1 cm (range, 0.5–2.5 cm). Complete removal of the US-visualized abnormality was achieved in 100% of patients. Ninety-five lesions (95%) had benign pathology. There were five lesions (5.0%) that were malignant and one atypia (1.6%), which was negative on routine pathologic review after re-excision open biopsy.

Hemostasis is achieved through the same tract with pressure most of the time (10 minutes), but sometimes the hematoma can be from injury to an artery resulting in surgical bleeding. Biopsy can result in a large expanding hematoma that causes high levels of anxiety for the patient and the surgeon. In the past, the only way to stop this surgical bleeding was by taking the patient to the operating room for exploration. We described a novel technique to control the bleeding utilizing a pediatric foley catheter to tamponade the vessel from inside and out (24).

In patients >40 years, a 6-month follow-up with mammography documented total resolution of the mammographic lesion, demonstrating that vacuum-assisted excisional breast biopsy under US guidance is an effective technique for therapeutic management of benign lesions, with minimal morbidity and optimal cosmesis. At present, we routinely remove benign-appearing lesions in this fashion (23).

Table 1  Comparison of vacuum assisted biopsy instruments

<table>
<thead>
<tr>
<th>Description</th>
<th>Mammotome®</th>
<th>VACU FLASH™</th>
<th>EnCor™</th>
<th>SUROS ATEC™</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single insertion</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>Upgradeable software</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
<td>N</td>
</tr>
<tr>
<td>Access to individual specimens</td>
<td>Y</td>
<td>Y</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Vacuum pulse on demand</td>
<td>Y</td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>MRI compatible</td>
<td>N</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
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Figure 4  Vacuum assisted biopsy with ultrasound guidance. (A) Ultrasound guidance is used to place the needle of the vacuum assisted device in place with the lesion of concern positioned within the biopsy sampling chamber; (B) the vacuum device can take multiple core biopsies with a single insertion of the needle, suctioning each sample into a collecting chamber that can be visualized by the operator throughout the procedure to evaluate for completeness. Note that the 8- or 11-gauge core samples are larger than the typical 12- or 14-gauge core needle biopsy samples.
Preparation for CNB or IVEB is carried out by the clinic staff. Local anesthesia will be accomplished with at least 5 cc of lidocaine HCL 1% with epinephrine 1:100,000 and 5 cc bupivacaine 0.5%. Patients who desire can be premedicated with oral valium. The breast lesion is removed with the Mammotome Breast Biopsy System® (Ethicon, Cincinnati, OH, USA) using US or mammographic guidance for placement of the 8- or 11-gauge probe. The vacuum-assisted stereotactic biopsy is commonly used at our institution to obtain diagnosis and has been reported to completely remove cancerous lesions (pathologically) nearly 30–48 percent of the time (25,26). Thus, IVEB by itself may not be considered a therapeutic procedure for cancer, even when the target lesion is completely removed.

We have extensive experience using IVEB, not only under stereotactic guidance, but also under US guidance for same-day excisional biopsy of benign lesions. Our experience and that of others has shown that when stereotactic IVEB is used with intention to only incisionally biopsy a suspicious lesion, the result is complete lesion removal in approximately 30–48% of cases (25,26). This indicates its potential utility for complete lesion removal of mammographically detected lesions and determination of size utilizing IVEB (20). Another disadvantage of using IVEB to both obtain a diagnosis and completely excise a lesion is the inability to reconstruct the margin status of a malignant tumor. Thus, after an IVEB procedure for diagnostic purposes, a malignant diagnosis requires lumpectomy for margins. Since our long-term goal is to develop a same-day diagnosis and first-line treatment regimen for small, unicentric breast cancers, we performed a pilot trial of percutaneously resect (IVEB)/percutaneous margin ablation/open resect protocol (23). In this way, using IVEB in combination with ablative techniques, we achieved oncologic excision of tissue for evaluation and negative margins by ablation with 100 percent success (23).

Alternative image-guided excisional techniques may remove the tumor intact. Presently available options are less well developed and were not chosen for this work secondary to significant concerns of tract seeding, cautery distortion of diagnosis (Neothermia™), and larger incision size [Advanced Breast Biopsy Device (ABBI™)] while still not accomplishing margin negativity or improved cosmesis (17).

US-core biopsy of axillary lymph nodes

The most important prognostic indicator involved in the evaluation of a patient with invasive breast cancer is axillary staging (27). This data is typically obtained in the operating room under general anesthesia with utilization of a nuclear medicine technetium injection in order to perform sentinel lymph node biopsy. This process can be a time consuming and costly undertaking requiring many healthcare resources. A variety of non-invasive imaging modalities in conjunction with biopsy techniques have been studied to evaluate the axilla for metastatic disease in an effort to comply with efficiency and cost savings standards during this era of healthcare reform. PET/CT has a sensitivity and specificity of identifying malignant nodes of 56 and 96 percent, respectively (28–30). Similar results were demonstrated with MRI, finding a sensitivity of 66% and specificity of 93% (31-33). Ultrasonography used for axillary staging has a sensitivity of 87% and specificity of 98%, and its use provides significant cost savings in comparison with both PET-CT and MRI (34). With ultrasonography being at least equal in accuracy and more cost efficient than the alternatives, this modality has been increasingly employed as a method to determine nodal positivity (34-36).

Suspicious nodes on US imaging can be specifically targeted for tissue confirmation through use of FNA or core biopsy or vacuum-assisted biopsy methods under local anesthesia, in turn saving the patient a trip to the operating room (37,38). Positive nodal tissue on pathology review is an indication for the patient to proceed to neoadjuvant chemotherapy. A negative node biopsy potentiates the decision to proceed with staging axillary sentinel node biopsy or definitive standard management based on breast cancer stage.

A retrospective review of patients at our institution from 2007–2012, included patients undergoing US of the axilla with CNB (39). A 95 patient cohort was divided into clinically positive (32%) and clinically negative (68%) axillae. The sensitivity and specificity of axillary US guided CNB was 90% (95% CI: 84.8% to 98.8%) and 100% (95% CI: 77% to 100%), respectively, and demonstrated cost savings of $485,007 compared to operative sentinel node biopsy. The axillary US guided CNB done under local anesthesia in the office setting can avoid the need for an additional operation, saving significant time and resources in patients with breast cancer who may have nodal involvement (39).

**Touch preparation cytology (TPC) on core biopsies**

TPC or imprint cytology has proven to be reliable and
more efficient than other methods of intraoperative evaluation for tumor diagnosis and margin evaluation (40,41). TPC is accomplished by touching or smearing the core biopsy or lesion to a glass slide. Tumor cells and benign cells will adhere to the slide, but fat cells will not. The slide is then fixed with 95% methanol and stained with hematoxylin and eosin. At our institution, we evaluated 428 patients for intraoperative diagnosis and margin evaluation by touch preparation analysis. There were 83 malignancies, of which 26.5% were in situ. The average tumor size was 2.2±1.9 cm. The diagnostic accuracy was 99.3%; sensitivity was 96.4%; and specificity was 100%. The positive predictive and negative predictive values were 100% and 99.3%, respectively (40). When compared to frozen sections, TPC has also proven to be more efficient and reliable, with lower false positive and false negative rates (41). We published equally accurate results for the intraoperative diagnosis of sentinel lymph nodes evaluated with TPC (42,43). We extended this concept to TPC of CNB specimens as a means of same-day diagnosis in the outpatient setting (44). A total of 55 outpatients presenting with breast lesions underwent 8- or 11-gauge vacuum-assisted core or 14-gauge biopsy gun CNB and TPC for diagnosis. In group I, CNB specimens were sent to pathology for TPC processing. In group II, TPC of the CNB specimen was performed immediately by the surgeon, and slides were then sent to pathology to be read. These results were compared to permanent pathology sections of the CNB. The sensitivity, specificity, positive predictive value, and negative predictive value of TPC in group I was 62.5%, 100%, 100%, and 83.3%, respectively, with a false negative rate of 37.5%. The low sensitivity of Group I was felt to be secondary to drying of the core specimens during transport to the pathologist. Also, in this group lobular cancer was not detected by TPC. When TPC was performed immediately on site (group II), both the sensitivity and specificity of TPC were 100%, with a zero false-negative rate.

Others have reported fair to good results in terms of accuracy in predicting benign versus malignant results (45). This may be due to drying of specimens before they reach pathology. We advocate a coordinated effort between the pathologist and the surgeon such that the surgeon prepares the slides immediately and fixes them in alcohol and then sends these slides to pathology for staining and cytologic interpretation. Alternatively, the cytologist can come to the biopsy suite.

Open excisional breast biopsy

Needle localization breast biopsy (NLBB) is the standard for removal of non-palpable breast lesions after vacuum-assisted core biopsy. Disadvantages include a miss rate of up to 22%, a positive margin rate of up to 75%, and vasovagal reactions (~20%) (46,47). Surgeons cannot rely on needle localization of the clip alone and must be cognizant of potential clip migration. We performed a retrospective review of post-biopsy films in patients who had undergone vacuum-assisted core biopsy with stereotactic clip placement for abnormal mammograms (48). We measured the distance between the clip and the biopsy site in standard two-view mammograms, using the Pythagorean theorem to calculate the distance the clip moved within the breast from the biopsy site. Pathology reports on lesions removed by NLBB versus those removed by intraoperative US-guided breast biopsy, which uses US to localize the core biopsy site created by vacuum excision, were reviewed to assess margin status. A total of 165 post-biopsy mammograms on patients who had vacuum-assisted core biopsy with clip placement were reviewed. In 93 evaluable cases, the mean distance the clip moved was 13.8±1.6 mm (SEM; 95% CI: 10.6–17.1 mm). The range of migration was 0–78.3 mm. The median was 9.7 mm. In 22.5% of patients, the clip was >20 mm from the targeted site. Migration of the clip was not related to the age of the patient, the size of the breast, or the location within the breast. In the subgroup of patients with cancer, margin positivity after NLBB was 42% versus 0% when the clip was ignored and the cavity removed by intraoperative US-guided biopsy (48). These results led us to develop the patented methodology used in the present protocol to locate the vacuum-assisted core biopsy cavity via US.

Hematoma or cavity-directed US-guided excisional biopsy after stereotactic vacuum-assisted breast biopsy

To decrease the miss rate that can occur with NLBB, we hypothesized that the hematoma-filled cavity resulting from a CNB vacuum-assisted core biopsy could be visualized with US and used to guide excision (Figures 5,6) (49,50). The hematoma-filled cavity visualized by US, and then grossly at surgery, could also be seen microscopically to again confirm lesion removal (49,50). In a study of 455 patients we compared hematoma-directed US-guided procedure for intraoperative localization of non-palpable lesions to needle localization done at the same institution.
Needle localization was performed on 126 (28%) patients and 329 (72%) via HUG. The previous core-biopsy site in 100% of patients was successfully excised using HUG: 152 of 329 (46%) were benign and 177 of 329 (54%) were malignant. Margins were positive in 42 of these 177 cases (24%). A total of 88 of 126 (70%) were benign and NLBB 38 of 126 (30%) were malignant; margins were positive in 18 of these 38 (47%) with NLBB. Margin positivity was significantly higher for NLBB than HUG (P=0.045, Fisher exact). This study provides proof of concept as to the effectiveness of US in identifying the cavity of the biopsy site. We have performed the procedure up to 5 weeks after stereotactic biopsy, which allows easier scheduling. In addition, it improves patient comfort over needle localization. We now routinely use hematoma-directed US guidance as a standard localization procedure in lieu of NLBB in this breast program and propose to use it for localization of the vacuum cavity for the ablative procedures.

**Summary**

In summary, US is an extremely valuable tool in breast surgery, with a variety of uses including screening, diagnostic imaging, image-guided biopsy techniques, image-guided percutaneous excision techniques, as well as image-guided open surgical excision procedures. Technology has advanced to include improved imaging and vacuum-assisted devices making percutaneous biopsies and excisions easier and more efficient. We have found touch prep cytology to be accurate and allows for rapid pathologic results of a tissue specimen, regardless of the source being from breast tissue or from a lymph node. This provides diagnostic capability of a breast lesion with potential staging of the axilla in the clinic setting with cytology results the same day and tissue for marker assessment. We believe this provides optimal care for the patient allowing for thorough pre-operative planning in order to minimize unnecessary additional operations. In the future, we look to improve upon this foundation of tools with implementation of percutaneous excision of lesions followed by ablation of the surrounding tissue for definitive complete treatment of early stage breast cancer in the clinic setting.

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None.

**Footnote**

Conflicts of Interest: The authors have no conflicts of interest to declare.

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