**Influence of Biospecimen Variables on Proteomic Biomarkers in Breast Cancer**


**Abstract**

**Background:** PI3K/Akt/mTOR signaling is being actively pursued as a therapeutic target for breast cancer. We sought to determine if tumor heterogeneity and biospecimen variables affect the evaluation of PI3K/Akt/mTOR pathway markers.

**Methods:** Intraoperative image-guided core-needle biopsies (CNB), and central and peripheral surgical tumor specimens were prospectively collected in 53 patients with invasive breast cancer. Specimens were assessed with reverse-phase protein arrays (RPPA) and immunohistochemistry (IHC).

**Results:** There was a moderate or strong correlation between the expression of 149 (97%) of the 154 different RPPA markers in the center and periphery. Correlation was higher for smaller tumors, in patients who did not undergo neoadjuvant therapy, and with shorter cold ischemia time. Of 154 markers, 132 (86%) were not statistically different between the center and periphery, and 97 (63%) were not different between the CNB and the surgical specimen (average of the central and peripheral specimen). pAkt S473 and PTEN had a significant correlation between central and peripheral specimens, and between CNB and surgical specimen. However, pAkt S473, pS6 S235/236, and pS6 240/244 levels were significantly higher in CNB than the central specimens both by RPPA and by IHC.

**Conclusions:** Most individual proteomic biomarkers studied do not have significant intratumoral heterogeneity. However, protein and phosphoprotein levels are affected by biospecimen type and other preanalytic variables. PI3K pathway activation is greater in CNB compared with postexcision surgical samples suggesting a potential loss of phosphorylation during surgical manipulation, or with cold ischemia of surgical specimens. Clin Cancer Res; 20(14); 3870–83. ©2014 AACR.
Translational Relevance

There is great interest in using protein-based biomarkers, especially phosphoproteins for assessment of pathway activation, as potential predictors of response to targeted therapies. Our results demonstrate that most individual proteomic biomarkers did not have significant intratumoral heterogeneity. However, protein and phosphoprotein levels were affected by biospecimen type and other preanalytic variables. PI3K pathway activation was greater in core-needle biopsies compared with surgical samples. Use of archival samples, especially surgical samples may introduce additional variability to biomarker assessment. These findings have implications for study design for biomarker-driven targeted therapy trials as well as window of opportunity studies. These studies also highlight the need for biospecimen science to reduce postexcision degradation of phosphorylation as well as the need for identifying the most robust markers least prone to intratumoral heterogeneity and biospecimen and assay-related variation.

We hypothesized that biospecimen variables and tumor heterogeneity may affect assessment of pathway activation. We therefore performed a prospective clinical study to determine whether biospecimen variables including specimen type (in vivo collection of FNA or core biopsy vs. ex vivo collection of central and peripheral surgical tumor samples) affect the functional proteomic profile by RPPA and the assessment of PI3K/Akt/mTOR pathway activation by RPPA and IHC. This work was performed to assess effect of biospecimen variables on RPPA results without microdissection. We demonstrated that most biomarkers do not have significant spatial heterogeneity in operable breast cancer. However, levels of several potentially actionable biomarkers, including markers of PI3K pathway activation, are affected by biospecimen type and biospecimens handling variables.

Materials and Methods

Human specimen collection

We conducted a prospective laboratory study after approval by the institutional review board (Fig. 1). After informed consent, 53 patients with invasive breast cancer 2 cm or greater on preoperative imaging underwent intraoperative ultrasound-guided percutaneous FNs (4–8 passes) and CNBs (3–4 core biopsies) after induction of anesthesia but before the beginning of the operation. The number of samples obtained per patient is listed in Supplementary Table S1. FNs were frozen for RPPA. In addition, one CNB with the greatest tumor content upon visual inspection was macrodissected and frozen for RPPA. The other CNBs were formalin fixed and paraffin embedded (FFPE). The surgical specimen was evaluated by a dedicated breast pathologist in the frozen section suite. For breast conservation (lumpectomy) samples, whole specimen radiographs were performed for confirmation of target lesion removal. For both breast conservation and mastectomy samples, margins were inked and sliced specimens were radiographed as deemed necessary for margin analysis as previously described (7). A portion of central and peripheral tumor was then allocated for research; one specimen from each site was snap frozen for RPPA, another was placed in formalin. Central and peripheral research samples were acquired at the same time, and the samples were approximately the same size, and were frozen directly. Small specimens of central and peripheral tumor samples were removed from main specimen in the frozen section suite and then FFPE directly. Surgical tumor samples were macrodissected, microdissection was not performed. The time from intraoperative specimen removal to time of freezing of the research specimen was prospectively recorded and designated as "cold ischemia time."

The FFPE samples were assessed by hematoxylin and eosin (H&E) staining. Three patients had no tumor on H&E staining of all 3 samples (CNB, central, and peripheral specimen). These patients were excluded from further analysis. Twelve patients had no tumor on H&E of 1 of the 3 samples. The samples without tumor were excluded from analysis.
Reverse-phase protein arrays

RPPA was performed in the MD Anderson Cancer Center Functional Proteomics Core Facility as described previously (10). Lysates were extracted from each tumor specimen and 4 replicates from each lysate were spotted on the RPPA. The cell lines were treated on separate days to obtain 3 biological replicates. Briefly, 200 μL of lysis buffer was added to CNB samples. Following homogenization with ceramic beads, samples were centrifuged, and supernatant was quantitated using BCA Assay Kit (Thermo Scientific). The final protein concentration was adjusted to 2 μg/mL. FNA samples were lysed in 75 μL of lysis buffer that was already mixed with 25 μL of loading buffer. Following sonication, samples were quantitated with BCA Assay Kit and the final concentration was adjusted to 3 μg/mL. Lysates were probed with antibodies validated for RPPA that are enriched for components of PI3K/Akt/mTOR pathway (Supplementary Table S2). A total of 154 unique proteins (including phosphoproteins) and 4 replicates were evaluated. Protein levels were expressed as the mean expression values in Log2. Samples with less than 20% tumor cellularity on corresponding H&E sections were not included in statistical analysis for RPPA data.

Immunohistochemistry

IHC was performed on FFPE CNBs, and central and peripheral tumor specimens with the following antibodies: pAkt S473 (Cell Signaling #4060, 1:50, 1 hour), pS6 S235/236 (Cell Signaling #4858, 1:50, overnight), pS6 S240/244 (Cell Signaling #2215, 1:200, overnight), p4EBP1 T70 (Cell Signaling #9455, 1:50, overnight) Ki67 and PTEN immunostaining was performed in the MD Anderson IHC Core laboratory with the following antibodies: Ki67 (Dako #M7240, 1:100, 20 minutes) and and PTEN (Dako #M3627, 1:100, 15 minutes). Positive and negative controls were included for all assays. H-score was determined by estimation of the percentage of tumor cells positively stained with weak, moderate, and strong staining intensity (by A. Sahin). The cytoplasmic score was used for statistical analysis. The following formula was used: H-score = \[(\text{low%})/C_1 + (\text{medium%})/C_2 + (\text{high%})/C_3\].

Statistical analysis

The RPPA spot signal intensity data from MicroVigene were processed using the R package SuperCurve (version 1.4.3; ref. 1), available at "http://bioinformatics.mdandon-son.org/OOMPA." RPPA superslide was normalized for FNA and surgical samples/CNB samples separately. Raw data were treated with median centering across samples; the treated data were undertaken with centering by the sample medians; and the final normalized data are obtained by applying MAD scaling to the data; the median centered values were divided by median absolute deviation to scale the data (11-13). For Spearman’s rank correlation test, P values are computed using algorithm Applied Statistics (AS) 89 (14, 15). The values of correlation were classified as follows: rs of 0.8 to 1, the correlation is very strong; between 0.6 and 0.79, correlation is strong; between 0.5 and 0.59, the correlation is moderate; between 0.3 and 0.49, the
correlation is moderate to low; between 0.16 and 0.29, correlation is weak to low; and below 0.16, the correlation is too low to be meaningful. Hierarchical Ward-linkage clustering was performed based on Spearman correlation coefficients. Paired t test was conducted to determine the difference among RPPA data of various biospecimen types, and Wilcoxon signed-rank test was applied to compare IHC data. The Benjamini–Hochberg procedure was used to control the false discovery rate (FDR). FDR of 0.05 was considered significant.

For each of the 178 samples, we computed a PI3K pathway score, which was the sum of the normalized values (median centering and MAD scaling) of the phosphoprotein levels of Akt, mTOR, 4E-BP1, S6K, and S6 (i.e., PI3K score = pS6 240/244 + pS6 S235/236 + pS6K T389 + pmTOR S2448 + p4E-BP1 S65 + p4E-BP1 T37/46 + pPRAS40 T246 + pAkt S473 + pAkt T308). We classified the PI3K pathway scores of each sample type (FNA, CNB, central and peripheral surgical tumor) into 2 groups; samples were considered activated if their PI3K scores were in the top quartile within their sample type. Cohen k tests were performed to identify the agreement between PI3K pathway activity class and the biospecimen type. The k agreement was characterized as values <0 as indicating no agreement, 0 to 0.20 as slight, 0.21 to 0.40 as fair, 0.41 to 0.60 as moderate, 0.61 to 0.80 as substantial, and 0.81 to 1 as almost perfect agreement (16).

PTEN and pAkt were defined as loss and low, respectively, if their IHC values were 0. χ² tests were applied to check the dependency between PI3K score and PTEN loss/normal, PI3K score, and pAkt high/low. The analyses were performed by using R (version 2.11.1; R Development Core Team, 2011; ref. 17). Supplementary Table S3 describes how each point of REMARK guidelines were addressed.

Results
Clinicopathologic characteristics
Clinical and pathologic characteristics of the patients are listed in Supplementary Table S4. The median pathologic tumor size was 2.8 cm. Most patients (77%) had hormone-receptor positive tumors. Eleven patients had received neoadjuvant chemotherapy; these patients had their research biopsies intraoperatively (i.e., after completion of the neoadjuvant chemotherapy), followed by surgical excision, similar to the surgery first cohort. Most patients underwent a mastectomy. The specimen cold ischemia time was an average of 37 minutes, and ranged from 8 to 157 minutes.

A median of 6 FNA (range, 3–8) and 2 core-needle biopsy (range, 0–3) samples were obtained from each patient. Protein yield from a single FNA sample was a mean of 1,635 µg (median yield 686 µg; range, 36–8,580 µg). The protein yield from a single core biopsy was a mean of 862.8 µg (median yield 871.4 µg; range, 242–1,594 µg). The tumor cellularity of each sample was assessed by H&E and distribution of nonmalignant components in the central specimen including contribution of epithelium, lymphocytes and endothelium, and fibroblasts was detailed (Supplementary Table S5). There was no statistically significant difference in tumor cellularity between the central and peripheral samples, but the tumor cellularity was lower in the CNBs compared with the central specimens (in 47 matched samples 54.8% vs. 61.2%; P = 0.0689), and significantly lower in CNB compared with peripheral samples (in 43 matched samples 52.7% vs. 60.7%; P = 0.0476). Tumors whose corresponding FFPE samples had less than 20% tumor cellularity or that only had ductal carcinoma in situ (DCIS) on FFPE were considered unevaluable for RPPA, except for studies comparisons on effect of tumor cellularity. Tumors whose IHC sections did not contain tumor were considered unevaluable for IHC. Seven core biopsies, 4 central specimens, and 4 peripheral specimens had less than 20% tumor cellularity on FFPE. Forty-four (88%) of 50 CNBs, 46 (90%) of 51 of central tumor specimens, and 43 (84%) of 51 peripheral tumor specimens were considered evaluable for RPPA and IHC.

Effect of biospecimen type on proteome markers
To determine the effect of biospecimen type on the functional proteomic profile, we performed unsupervised hierarchical clustering on the RPPA from FNAs, CNBs, and central and peripheral surgical samples. As demonstrated in the heatmap (Fig. 2A), most of the FNA samples clustered together, whereas the central and peripheral samples and CNB samples coclustered suggesting that the cellular content of the FNA is markedly different from that of the CNB and surgical samples. Notably, CNB, central, and peripheral samples from the same patient often clustered together, consistent with intertumoral differences being greater than intratumoral differences. FNAs may not only differ because they are collected before any cold ischemia time, but also because there are previously reported differences in relative tumor vs. stromal cell content (18), and technical differences in RPPA sample processing and protein quantitation between FNA samples versus CNB and surgical samples. The correlation between different biospecimens for each biomarker is listed in Supplementary Table S6 and depicted as a heatmap (Fig. 2B).

There was a moderate or strong correlation between the central and peripheral specimen results for all but 5 of 154 proteins and phosphoproteins tested (Supplementary Table S6). There was a very strong correlation between the central and peripheral samples for ER (r = 0.896), PR (r = 0.90), and strong correlation for HER2 (r = 0.786; Fig. 3). We also compared the central and peripheral protein expression using paired Student t test (Supplementary Table S7). In 132 (86%) of 154 markers tested, at a FDR of 0.05, there was no statistical difference in the expression between central and peripheral specimens.

To determine the impact of marker assessment on in vivo biopsies compared with ex vivo excision samples, we next assessed whether RPPA results correlated between CNB and surgical specimens. There was a very strong correlation between the CNB and surgical tumor specimen (average of center and periphery) for ER (r = 0.804), PR (r = 0.871), and strong correlation with HER2 (r = 0.66; Fig. 3).
our data suggest that for especially ER, PR, quantitation by RPPA does not show significant differences attributable to spatial intratumoral heterogeneity or biospecimen type. There was no statistically significant difference in expression of 97 (67%) of the 154 markers assessed in CNB, compared with the surgical samples by Student $t$ test, at a FDR of 0.05 (Supplementary Table S7).

There was a modest to poor correlation between FNA results and surgical specimens for several proteins (Supplementary Table S6). However, the FNA and surgical sample had a significant correlation for 49 proteins, and the FNA and CNB had a correlation for 68 proteins. There was strong correlation between FNA and surgical samples for ER ($r = 0.606$), PR ($r = 0.695$), and fair correlation for HER2 ($r = 0.367$). There was also a strong correlation between CNB and FNA results for ER ($r = 0.650$), PR ($r = 0.642$), and moderate correlation for HER2 ($r = 0.57$).

**Effect of biospecimen variables on biomarker expression**

We next evaluated the effect of preanalytic biospecimen variables on biomarker expression. At FDR < 0.05, in FNA samples there was no difference in protein expression between patients that were younger versus older than median age of 51 years. In CNB, ER was higher in older patients ($P < 0.0001$), and TIGAR and c-Kit were higher in younger patients ($P = 0.0008$ and 0.001, respectively). ER expression also was significantly higher in central and peripheral samples in patients that were older ($P < 0.0001$ for both).

There was an increased pGSK3a/b S21/9 expression in grade 3 tumors compared with grade 1 or grade 2 tumors ($P = 0.0343$ and 0.0001, respectively). There was no protein differentially expressed between tumors of different stages, or between small versus large tumor size (median cut off 2.8 cm) at FDR < 0.05 cut-off for any of the FNA, core, central, and peripheral samples.

We next determined whether preanalytic biospecimen variables impacted the correlation of expression between different biospecimen types. We expected that larger tumors would have greater tumor heterogeneity, and indeed by paired $t$ test, we found that correlation between central and peripheral specimens was better in tumors that were smaller than the mean ($<3.6$ cm), compared with tumors larger than the mean ($P < 0.001$; Supplementary Table S8). Interestingly, larger tumors had a greater correlation between FNA versus surgical samples ($P = 0.003$). Tumors with shorter than the mean cold ischemia time (36.7 minutes) had better correlation between central versus peripheral specimens compared with tumors with longer cold ischemia time ($P < 0.001$; Supplementary Table S8). There was also better correlation between FNA and central specimens in tumors with shorter cold ischemia time ($P < 0.001$). Shorter cold ischemia time also was associated with a higher correlation between CNB and surgical specimens ($P < 0.001$).

We also assessed the effect of surgery type on biomarker correlation. There was a stronger correlation between CNB...
versus surgical samples ($P < 0.001$) as well as FNA vs. surgical samples ($P = 0.005$) in patients who had breast conservation compared with those who underwent mastectomy. There was also a trend toward improved correlation between central and peripheral samples in breast conservation samples ($P = 0.074$). To better understand whether other preanalytic variables differed between breast conservation and mastectomy samples, we compared tumor size and cold ischemia time for these specimens. There was no statistically significant difference in tumor size between patients who underwent breast conservation versus mastectomy (median size 3.1 cm vs. 3.7 cm; $P = 0.49$). However, surgery type was associated with differences in cold ischemia time. Mean cold ischemia time for breast conservation patients was 28.5 minutes (range, 20–42 min), compared with 39.1 minutes (range, 8–157 min) for mastectomy ($P = 0.021$). This may be attributable to the fact that breast conservation samples routinely undergo immediate margin analysis or maybe because mastectomy specimens take longer to section.
The correlation between FNA and surgical specimens was significantly greater for nonphosphorylated proteins than phosphorylated proteins \((P = 0.043)\). There was also a trend for difference in correlation between CNB and surgical specimens \((P = 0.055)\). Thus, phosphoprotein levels are more susceptible to biospecimen type/sample handling than total proteins.

We next evaluated the effect of tumor cellularity on biomarker expression (Supplementary Table S9). At FDR \(< 0.05\), there are 54 proteins in CNB data showing significant variability based on tumor cellularity. At FDR \(< 0.05\), there were 22 proteins in central tumor data showing significant variability based on tumor cellularity. There were 7 common proteins in CNB and central data that varied by tumor cellularity (cyclin B1, Beclin, IRS, YB1, p38 MAPK, BID, and eIF4G). The contribution of the tumor cell composition was then correlated with the variation in the expression of each marker between biospecimens. \(F\) test in linear regression model was applied to check if the tumor cell composition affects the variance in the expression of each marker. The variance of most of the biomarkers could not be explained by the tumor cell composition.

The biomarker correlation in the patients who did and did not receive neoadjuvant chemotherapy was also analyzed separately (Supplementary Table S10). Correlations for samples from patients who did not receive neoadjuvant chemotherapy were significantly higher than in patients who received neoadjuvant chemotherapy when the comparing the central versus peripheral specimen \((P < 0.0001)\), CNB versus central specimen \((P < 0.0001)\), CNB versus surgical specimen \((P < 0.0001)\), FNA versus CNB \((P < 0.0001)\), and FNA versus peripheral sample \((P = 0.0209)\); Supplementary Table S11).

### Effect of biospecimen type on PI3K/Akt/mTOR pathway activation

As the PI3K/Akt/mTOR pathway is a potential therapeutic target under active clinical investigation, we assessed PI3K/Akt/mTOR pathway activation by specimen type. First, we assessed expression of pAkt S473 and pAkt T308, markers of activated Akt (Fig. 3). pAkt S473 as measured by RPPA was correlated between all specimens and pAkt T308 was correlated between the central versus peripheral and core versus surgical specimen.

PTEN and INPP4B are 2 tumor suppressors whose loss is associated with activation of PI3K/Akt/mTOR signaling (19). The PTEN expression of central and peripheral samples and CNB versus surgical samples were strongly correlated, both \(r = 0.716\). The INPP4B expression of central and peripheral samples and CNB versus surgical samples were strongly correlated, both \(r = 0.774\).

Phosphorylation of mTOR target p4E-BP1 T37/46 correlated strongly between central and peripheral samples \((r = 0.615)\), whereas p4E-BP1 S65 correlated moderately \((r = 0.531)\); Fig. 3). p4E-BP1 T37/46 and p4E-BP1 S65 had moderate-to-low correlation between the CNB and surgical specimen \((r = 0.349 \text{ and } 0.318, \text{ respectively})\). There was also a moderate-to-low correlation between central and peripheral samples for the mTOR downstream signaling markers pS6 S235/236 and pS6 240/244 \((r = 0.416 \text{ and } 0.439)\). The CNB and surgical specimens had no meaningful correlation between CNB and surgical specimen for pS6 S235/236 \((r = 0.098)\) and weak-to-low correlation for pS6 240/244 \((r = 0.25)\). The correlation between the CNB and FNA was somewhat stronger for pS6 S235/236 \((r = 0.364)\) and pS6 240/244 \((r = 0.415)\).

As the concordance of PI3K pathway activation in the CNB and surgical specimens was moderate to low, we next determined whether there was a significant difference in PI3K pathway activation in CNB versus surgical specimen by paired Student \(t\) test. When the CNB was compared with the central specimen, expression was higher in CNB for pAkt S473 \((P = 0.020)\), p4E-BP1 S65 \((P = 0.025)\), p4E-BP1 S37/46 \((P < 0.001)\), pS6 S235/236 \((P = 0.011)\), and pS6 S240/244 \((P < 0.001)\), as well as PTEN levels \((P < 0.001)\) and pMAPK T202/204 \((P = 0.004)\; \text{Fig. 4})

We compared with the overall surgical specimen (average of the center and periphery), p4E-BP1 T37/46 and pS6 240/244 levels in the CNB were significantly higher \((P = 0.005\) and \(P < 0.001\), respectively).

### PI3K/Akt/mTOR signaling by IHC

We determined PI3K/Akt/mTOR signaling using IHC. Examples of positive and negative staining for each antibody are shown in Fig. 5A.

There was a very strong correlation in p4E-BP1 T70 expression in the central and peripheral tumor \((r = 0.81)\). There was a moderate correlation in central and peripheral expression of pAkt S473 \((i = 0.588)\), pPRAS40 T246 \((r = 0.82)\), pS6 S240/244 \((r = 0.553)\), and moderate-to-low correlation for pS6 S235/236 \((r = 0.475)\; \text{Fig. 5B})

There was a very strong correlation in PTEN expression in the central and peripheral tumor \((r = 0.85; \text{Fig. 5B})\). When PTEN IHC was categorically classified as having PTEN loss versus not, the central and peripheral samples had almost perfect agreement \((k = 0.916)\). CNB and central specimen had substantial agreement \((k = 0.682)\), and CNB and peripheral specimen had moderate agreement \((k = 0.579)\).

The CNB and central specimen immunostaining also correlated for pAkt S473 \((r = 0.87)\), PTEN \((r = 0.72)\), p4E-BP1 T70 \((r = 0.81)\), and pS6 S235/236 \((r = 0.38)\), but not for pPRAS40 T246 or pS6 S240/244. The CNB and peripheral specimen immunostaining were correlated for pAkt S473 \((r = 0.71)\), PTEN \((r = 0.68)\), p4E-BP1 S65 \((r = 0.63)\), pS6 S240/244 \((r = 0.47)\), and pS6 S235/236 \((r = 0.49)\).

We next compared PI3K pathway activation status by IHC to that of RPPA. pAkt S473 quantitation by RPPA correlated with immunostaining by IHC in central specimen \((r = 0.71)\), peripheral specimen \((r = 0.63)\), and CNB \((r = 0.52; \text{Fig. 5C})\). RPPA quantitation of PTEN correlated with IHC results in the central specimen \((r = 0.41)\), but not in the peripheral tumor or CNB specimens. pS6 S235/236 expression by RPPA and IHC also were significantly correlated in the central specimen and in the CNB, but not tumor periphery \((r = 0.24)\). When RPPA quantitation of FNAs was
compared with IHC of central specimens, there was moderate correlation for pAkt S473 \( (r = 0.30) \), PTEN \( (r = 0.37) \), and pS6 S235/236 \( (r = 0.35) \).

There was a very strong correlation between percentage cells positive for Ki67 between the central and peripheral specimen \( (r = 0.865) \) and between the CNB and central specimen \( (r = 0.891) \). There was no significant correlation between the expression of proliferation marker proliferating cell nuclear antigen (PCNA) on RPPA and percentage cells positive for Ki67 on IHC.

We then determined whether there were differences in immunostaining by biospecimen type. The pS6 S240/244 and pS6 S235/236 H-scores on the CNB were significantly higher than both the central specimen and the peripheral specimen \( (P < 0.001 \ for \ both; \ Fig. \ 6A) \). Examples of discordant IHC results with higher immunostaining in core biopsy than surgical specimens are shown in Fig. 6B.

**Concordance of PI3K pathway classification by RPPA**

We assessed pAkt S473, pAkt T308, pPRAS40 T246, p-mTOR S2448, p4E-BP1 S65, p4E-BP T37/46, pS6K T389, pS6 240/244, and pS6 S235/236 levels as indicators of high PI3K pathway activity, as they are known downstream targets \( (10, 20) \), and as their phosphorylation was found to be strongly inhibited by PI3K/mTOR inhibitor BEZ235 (100 nM) and Akt inhibitor MK-2206 (10 \( \mu \)M) in a panel of cancer cell lines \( (Fig. \ 7A \ and \ B) \).

When all samples were analyzed together, as expected, tumors with expression of pAkt S473 on IHC also had a significantly higher PI3K score than those that did not \( (P = 0.001; \ Fig. \ 7C) \). Furthermore, samples with PTEN loss by IHC \( (H\text{-score} = 0) \) had a significantly higher PI3K pathway activation score \( (P = 0.018; \ Fig. \ 7C) \). The statistical difference in PI3K activity score was not preserved when the threshold for PTEN loss was elevated to \( H\text{-score} = 50 \) or 100, suggesting that complete PTEN loss establishes the greatest differentiation between tumors that have PI3K activation versus not. When PTEN IHC and PI3K activity on RPPA was analyzed separately for each sample type (core, central, and peripheral), PI3K activity score was significantly higher in CNB that had PTEN loss versus those without PTEN loss by IHC \( (P = 0.0104) \), whereas there was no difference in PI3K activity score central or peripheral samples that had PTEN loss versus those without PTEN loss by IHC \( (Fig. \ 7D) \).

We then classified tumors as "PI3K signaling activated" or "not activated" based on PI3K activity score calculated separately for each specimen type. *F* test in linear regression model demonstrated that the variance of PI3K pathway score could not be explained by the tumor cell composition. The samples in the top quartile of PI3K RPPA core within each specimen type were classified as "PI3K signaling activated." In the 37 patients where RPPA results were available on both CNB and central specimens, the overall agreement between the core biopsy and central specimen was 78.4% \( (k = 0.454, P = 0.005) \). In the 39 patients for whom both central and peripheral RPPA results were available, the overall agreement was 74.4% \( (k = 0.33; P = 0.04) \). For samples that had all 4 biospecimen types were available, activity of PI3K pathway for each RPPA biospecimen type is shown in Fig. 7E.

**Discussion**

To deliver on the promise of personalized therapy, it is critical to develop robust markers that can assess activation of druggable targets such activation of PI3K/Akt/mTOR signaling in patient samples. Furthermore, in early clinical trials with targeted therapies, it is critical to incorporate...
pharmacodynamic studies to determine whether a new drug inhibits its target and by how much. Such studies rely on the ability to accurately assess proteomic markers in tumor samples. Marker assessment for patient selection is often performed on archival surgical tissue in FFPE. However, there are significant concerns about tumor heterogeneity and marker stability, especially for phosphoproteins. Therefore, we sought to determine how tumor heterogeneity and biospecimen variables affect assessment of proteomic biomarkers and, in particular, PI3K/Akt/mTOR pathway activation. Proteomic levels were differentially affected by biospecimen type and other preanalytic variables. For example, PI3K pathway activation was greater in CNB compared with surgical samples suggesting loss of phosphorylation during tumor handling during surgery and cold ischemia after removal from the patient.

In genomic studies, intratumoral heterogeneity was observed for mutations in oncogenes and multiple tumor-suppressor genes, even with gene-expression signatures of both good and poor prognosis being detected in different regions of the same tumor (6). Thus, we were concerned that there may be significant heterogeneity in

Figure 5. IHC for PI3K markers. A, IHC for PTEN, pAkt S473, pPRAS40 T246, p4E-BP1 S65, pS6 S240/244, and Ki-67. Examples of positive (top) and negative (lower) immunostaining for each marker are included. B, correlation of immunostaining for PTEN, pAkt S473, pS6 S240/244, and pS6 S235/236 between central and peripheral specimens (left) and central specimens and core-needle biopsy (right). Immunostaining was quantitated by H-score. C, correlation of central RPPA results with core-needle biopsy specimen immunostaining for PTEN, pAkt S473, pS6 S240/244, and pS6 S235/236.
Thus, in less cellular tumors PTEN loss may not be as
detectable as in more cellular tumors. PTEN expression in stroma as well as tumor cells.
Dissection, quantitation of PTEN in the tumor lysate represents a positive control in IHC. On RPPA, in the absence of microdissection, quantitation of PTEN in the tumor lysate represents PTEN expression in stroma as well as tumor cells. Thus, in less cellular tumors PTEN loss may not be as apparent on RPPA, and this may contribute to weak correlation between IHC and RPPA. PTEN IHC is attractive as not only does it seem to have low interspecimen variability, but also PTEN loss and in particular complete loss \( (H\text{-score} = 0) \) on IHC was associated with PI3K pathway activation by RPPA. Larger studies are needed to further test the impact on "low" PTEN on pathway activation in order to set the appropriate threshold for PTEN loss that is biologically and clinically meaningful.

There has been increasing concerns that specimen type (such as CNB vs. surgical sample) may affect biomarkers. When standard breast cancer markers in CNB were compared with excisions, discordance between CNB and excision for HER2 and ER expression was found to be rare (1.8% and 1.2%, respectively), however discordance in PR expression was common (15%; ref. 21). A recent meta-analysis that included 21 articles involving nearly 2,500 patients compared concordance of ER and PR in CNBs and excision and found an overall agreement between CNB and surgical excisions of 92.8% for ER \( (k = 0.78) \) and 85.2% for PR \( (k = 0.66) \); ref. 22). In contrast, Baker and colleagues reported that, in colorectal cancer, pAkt S473 staining was observed only in biopsies and not in surgical samples (albeit not assessed with RPPA; ref. 2), thus authors suggested that caution should be used when using phosphoprotein levels in surgical specimens to measure intrinsic signaling activity or drug effects because of the potential for rapid dephosphorylation. Recently, to evaluate the impact of ischemia, Mertins and colleagues collected ovarian tumor and breast cancer xenograft and performed quantitative proteomics and phosphoproteomics with RPPA and liquid chromatography/mass spectrometry after defined ischemic intervals (23). Although the global expressed proteome and most of the >25,000 quantified phosphosites were unchanged after 60 minutes, rapid phosphorylation changes were observed in up to 24% of the phosphoproteome. Concern about stability of cell signaling markers was also raised in a recent study on breast cancer, in which loss of pAkt and pErk was found by IHC on CNB obtained ex vivo but not in surgical samples.
after routine processing (3). These studies suggest that loss of phospho-staining can occur during routine fixation of resection specimens. Our study design was different than the latter breast cancer study as our CNB were obtained not on surgical samples ex vivo, but in the operating room in vivo, while samples of surgical specimens were frozen or formalin fixed immediately upon specimen processing for clinical care, thus mimicking earliest potential time point for specimen collection for tumor banking or biomarker assessment. Our study findings are consistent with the previous reports in that pAkt and pS6 levels were higher in CNB, and there were several tumors in which the PI3K score was elevated in the CNB, but not in the surgical sample. Taken together, our results suggest that PI3K pathway activation is greater in vivo, with subsequent dephosphorylation/pathway inactivation ex vivo.

PI3K/Akt/mTOR pathway activity is being pursued as a potential predictor of response, marker of target inhibition,
suggesting that these variables may contribute to the dynamic range and total observable markers. Indeed, ER, PR, and HER2 all tend to have a large FNA and surgical sample as well, including standard of proteomic markers and to assess pharmacodynamics demonstrate the prognostic role of known and novel features that several studies have successfully used FNAs to demonstrate the prognostic role of known and novel proteomic markers and to assess pharmacodynamics markers of response to targeted therapies (10, 25).

In summary, overall there is limited spatial intratumoral heterogeneity for individual proteomic markers in operable breast cancer. However, some actionable markers are differentially expressed by biospecimen type and other preanalytic variables. Although expression of most markers on CNB and surgical samples correlate, some do differ. Notably, PI3K pathway activation is greater in CNB compared with surgical samples, thus CNB may be preferable to surgical samples to assess pathway activity. Careful study of preanalytic and analytic variables that influence results is needed for clinical implementation of each biomarker.
Disclosure of Potential Conflicts of Interest

G. B. Mills reports receiving commercial research grants from AstraZeneca, Celgene, CelStines, Exelixis/Sanoﬁ Aventis, GlaxoSmithKline, Roche, SDI, and Wyeth/Pfizer/Puma; has ownership interests in Catena Pharmaceuticals, PTV Ventures, Spindle Top Ventures; and is a consultant/advisory board member for Arxis Biotechnology, Acuragen, Aushon, Catena Pharmaceuticals, Daiichi Pharmaceuticals, Foundation Medicine, Han Allio Korea, Komen Foundation, Novartis, Targeted Molecular Diagnostics, and Tau Therapeutics. No potential conﬂicts of interest were disclosed by the other authors.

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