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Analytic Validation of a Clinical-Grade PTEN Immunohistochemistry Assay in Prostate Cancer by Comparison to *PTEN* FISH

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Abstract

PTEN loss is a promising prognostic and predictive biomarker in prostate cancer. Because it occurs most commonly via *PTEN* gene deletion, we developed a clinical-grade, automated and inexpensive immunohistochemical assay to detect PTEN loss. We studied the sensitivity and specificity of PTEN immunohistochemistry relative to 4-color fluorescence in situ hybridization (FISH) for detection of PTEN gene deletion in a multi-institutional cohort of 731 primary prostate tumors. Intact PTEN immunostaining was 91% specific for absence of PTEN gene deletion, (549/602 tumors with 2 copies of the PTEN gene by FISH showed intact expression of PTEN by immunohistochemistry) and 97% sensitive for presence of homozygous PTEN gene deletion (absent PTEN protein expression by immunohistochemistry in 65/67 tumors with homozygous deletion). PTEN immunohistochemistry was 65% sensitive for presence of hemizygous PTEN gene deletion, with protein loss in 40/62 hemizygous tumors. We reviewed the 53 cases where immunohistochemistry showed PTEN protein loss and FISH showed 2 intact copies of the PTEN gene. On re-review, there was ambiguous immunohistochemistry loss in 6% (3/53) and failure to analyze the same tumor area by both methods in 34% (18/53). Of the remaining discordant cases, 41% (13/32) revealed hemizygous (n=8) or homozygous (n=5) PTEN gene deletion that was focal in most cases (11/13). The remaining 19 cases had 2 copies of the PTEN gene by FISH, representing truly discordant cases. Our automated PTEN immunohistochemistry assay is a sensitive method for detection of homozygous PTEN gene deletions. Immunohistochemistry screening is particularly useful to identify cases with heterogeneous *PTEN* gene deletion in a subset of tumor glands. Mutations, small insertions or deletions and/or epigenetic or microRNAmediated mechanisms may lead to PTEN protein loss in tumors with normal or hemizygous PTEN gene copy number.

Keywords

Prostatic carcinoma PTEN; fluorescence in situ hybridization; immunohistochemistry; radical prostatectomy; biomarker

Introduction

PTEN is the most commonly lost tumor suppressor gene in prostate cancer (1-5) and is a promising prognostic biomarker for poor clinical outcomes (6-18). Since the *PTEN* gene is almost always lost by genomic deletion of the entire gene in prostate tumors, fluorescence *in situ* hybridization (FISH) has traditionally been the gold standard assay to detect *in situ PTEN* loss in tumor tissue. However the relatively recent availability of reliable rabbit monoclonal antibodies for detection of PTEN protein has enabled the development of highly validated immunohistochemistry protocols to detect PTEN loss in prostate cancer (9, 19). Immunohistochemistry-based detection of PTEN loss in prostate cancer is less expensive and less time-consuming than FISH for the routine screening of prostate tumor specimens, making it easier to adapt to the current pathology work flow for risk assessment in prostate cancer. In addition, since *PTEN* loss is commonly subclonal and heterogeneous in primary prostate tumors (9, 20-22), detection of *PTEN* gene deletion by FISH can be technically challenging in some cases and screening for focal loss may be more easily accomplished by immunohistochemistry. Finally, there is emerging evidence that in addition to genetic

deletion, PTEN protein levels may be compromised by mutations in the gene or microRNAor epigenetic-regulated mechanisms which would not be detectable by FISH (9, 23-25).

We previously optimized and validated a PTEN immunohistochemistry assay for the detection of PTEN loss in prostate cancer specimens (9), and PTEN loss by this assay correlated with increased risk of biochemical recurrence in a case-control cohort of patients undergoing radical prostatectomy (12) and with risk of progression and metastasis in two high risk surgical cohorts (though the latter was not significant in multivariate analyses) (9, 11). Though originally performed manually, we have recently transferred this assay to a clinical-grade automated immunostaining platform that may be run in any CLIA-certified pathology laboratory. Using this assay, we recently reported that PTEN loss is associated with reduced recurrence free survival in multivariable models in a multi-institutional cohort of surgically treated patients (26) and with higher risk of lethal prostate cancer in a large population-based cohort (18). PTEN gene deletion by FISH has also been recently reported in a subset of the multi-institutional cohort and correlated with recurrence free survival (17). Here, to analytically validate our clinical-grade PTEN immunohistochemistry assay, we compared the performance of the automated immunohistochemistry assay to PTENFISH in this cohort, one of the largest multi-institutional cohorts to be studied by both techniques. We demonstrate that our immunohistochemistry assay shows robust sensitivity and specificity for detection of homozygous PTEN gene deletion.

Methods

Subject selection and tissue microarray design

The Canary Foundation Retrospective Prostate Tissue Microarray Resource has been described in detail elsewhere (27). Briefly, it is a multicenter, retrospective prostate cancer tissue microarray created as a collaborative effort with radical prostatectomy tissue from six academic medical centers: Stanford University, University of California San Francisco, University of British Columbia, University of Washington (including tissues from University of Washington and a separate cohort from the Fred Hutchinson Cancer Research Center), University of Texas Health Science Center at San Antonio, and Eastern Virginia Medical School. Tumor tissue from 1275 patients was selected for the tissue microarray using a quota sampling plan, from radical prostatectomy specimens collected between 1995 and 2004. A starting date of 1995 was selected to enrich for cases occurring after the implementation of PSA screening. There was no central pathology review in this cohort. The tissue microarray included samples from men with (a) recurrent prostate cancer; (b) nonrecurrent prostate cancer; and (c) unknown outcome due to inadequate follow-up time (ie, censoring). Recurrent cases of Gleason score 3+3=6 and 3+4=7 were relatively over-sampled as well as non-recurrent cases with Gleason score 4+4=8, in order to improve power to detect biomarkers providing prognostic information independent of Gleason score.

Each site built 5 tissue microarrays, each containing tumor tissue from 42 patients (210 patients from each contributing site). Each tumor was sampled in triplicate, utilizing 1 mm cores and an additional core of histologically benign peripheral zone tissue was included for each patient as a control. Recurrent and non-recurrent patients were distributed randomly across all tissue microarrays.

Immunohistochemistry assays

PTEN immunohistochemistry was performed on the CFRPTMR cohort as recently reported (26). Briefly, the protocol uses the Ventana automated staining platform (Ventana Discovery Ultra, Ventana Medical Systems, Tucson, AZ) and a rabbit anti-human PTEN antibody (Clone D4.3 XP; Cell Signaling Technologies, Danvers, MA). We previously validated a manual version of this assay using the same antibody in genetically characterized cell lines and prostate tumor tissue, showing strong correlation of the immunohistochemistry with *PTEN* gene copy number by 2-color FISH and high resolution SNP array analysis (9) and good correlation with 4-color FISH in a small cohort of needle biopsy specimens (28). To prove equivalence between the manual and automated assays, we also examined a test tissue microarray containing 50 prostate cancer cases with known PTEN protein status (including more than 30 with PTEN protein loss) by manual staining and found 100% concordance between the PTEN protein status on the manual and automated platforms.

Immunohistochemistry scoring

After staining for PTEN, all tissue microarrays were scanned at $20 \times$ magnification (Aperio, Leica Microsystems, Buffalo Grove, IL) and segmented into TMAJ for scoring (http://tmaj.pathology.jhmi.edu/). PTEN protein status was blindly and independently scored by two trained pathologists (TLL and CLM) using a previously validated scoring system (see below). Overall, there was "very good" agreement between independent reviewers, with 96% agreement over 2783 cores scored by both reviewers ($\kappa = 0.905$; 95% CI=0.887-0.923) (26).

A tissue core was considered to have PTEN protein loss if the intensity of cytoplasmic and nuclear staining was markedly decreased or entirely negative across >10% of tumor cells compared to surrounding benign glands and/or stroma, which provide internal positive controls for PTEN protein expression (9). If the tumor core showed PTEN protein expressed in >90% of sampled tumor glands, the tumor was scored as PTEN intact. If PTEN was lost in <100% of the tumor cells sampled in a given core, the core was annotated as showing heterogeneous PTEN loss in some, but not all, cancer glands (focal loss). Alternatively, if the core showed PTEN loss in 100% of sampled tumor glands, the core was annotated as showing homogeneous PTEN loss. Finally, a small percentage of cores were scored as having ambiguous PTEN immunohistochemistry results. This occurred when the intensity of the tumor cell staining was light or absent in the absence of evaluable internal benign glands or stromal staining. The percent of tissue cores with ambiguous scoring for PTEN immunohistochemistry was fairly constant across 6 of the 7 institutions included in the Canary tissue microarray cohort and varied from 0.7%-5.3% (26).

For statistical analysis, each patient's tumor sample was scored for the presence or absence of PTEN loss by summarizing the scores of each individual sampled core from that tumor. A patient's tumor was designated as having heterogeneous PTEN loss if at least one tumor core showed heterogeneous PTEN loss, or alternatively, if at least one core showed heterogeneous or homogeneous PTEN loss and at least one core showed PTEN intact in tumor cells. A patient's tumor was scored as showing homogeneous PTEN loss if all sampled tumor cores showed homogeneous PTEN loss. Finally, a patient's tumor was scored

as having PTEN intact if all sample tumor cores showed intact PTEN in sampled tumor glands.

Initial blinded analysis of PTEN FISH

PTEN FISH was performed as previously described for a subset of this cohort (17). Briefly, the PTEN Del TECT FISH utilizes a four color probe combination as described. Probes were supplied by CymoGenDx LLC (New Windsor, NY) as follows: centromeric copy control probe - CYMO-Pink; *WAPAL* – CYMO-Green; *PTEN* – CYMO-Red; and *FAS* – CYMO-Aqua. We have shown previously that use of the probes bracketing *PTEN* improves the fidelity of assessments of PTEN loss (29). The two probes *WAPAL* and *FAS* on either side of PTEN provide information about the size of larger deletions and also allow recognition of background artifactual losses of *PTEN* due to histologic sectioning. Artifacts in assessing *PTEN* locus in cells in the section while leaving the centromere in place. The latter is a result of the long distance between the centromere and the *PTEN* locus on chromosome 10.

*PTEN*FISH analysis was performed entirely independently of PTEN immunohistochemistry, using 5 micron tissue microarray sections stained with DAPI (4',6diamidino-2-phenylindole, dihydrochloride) in tumor areas selected by a pathologist who was not involved in PTEN immunohistochemistry scoring (TJ) using an immediately adjacent section stained with hematoxylin and eosin. *PTEN* copy number was evaluated by counting spots for all four probes using SemRock filters appropriate for the excitation and emission spectra of each dye in 50–100 non-overlapping, intact, interphase nuclei per tumor tissue microarray core. For the initial blinded analysis of each case, two tumor-containing cores were scored based on the overall quality of FISH hybridization. In cases where different clonal deletions were present, all three cores were analyzed and more cells were analyzed. Hemizygous (single copy) *PTEN*loss was assigned when >50% of nuclei exhibited either interstitial loss of *PTEN* or concomitant loss of adjacent genes (*PTEN* and *WAPAL* and/or *FAS*). Homozygous deletion was defined by a simultaneous lack of both *PTEN*locus signals in 30% of scored nuclei.

Immunohistochemistry-guided re-analysis of cases with discrepant results by immunohistochemistry and FISH

53 cases showed PTEN protein loss by immunohistochemistry with 2 copies of *PTEN* gene present by initial FISH analysis (see Results, below). Two cases showed PTEN protein intact by immunohistochemistry with homozygous *PTEN* deletion by *PTEN* FISH. To analyze the cause of these discrepancies, we re-examined both the immunohistochemistry and FISH data in these cases. A digitally scanned photomicrograph of the most representative core with immunohistochemistry loss was selected to guide FISH re-analysis of the identical core from each case. Since the majority (85%) of these discrepant cases showed only focal immunohistochemistry loss in a subset of glands, the FISH re-analyses concentrated on determining the *PTEN* gene copy number within these small areas guided by the immunohistochemistry staining. Since only 50-100 cells from the best two of the three tumor-containing cores were initially analyzed for each case by *PTEN*FISH (29), this more

extensive analysis could include tissue microarray cores and regions of tissue microarray sections that had not been studied by FISH during initial blinded analysis.

Immunohistochemistry and FISH on standard tissue sections

To examine possible effects of tumor heterogeneity on immunohistochemistry and FISH interpretation in the setting of tissue microarray cores, we additionally examined 20 cases of varying PTEN status (enriched for discordance between immunohistochemistry and FISH) by FISH and immunohistochemistry on standard tissue sections. Immunohistochemistry and FISH interpretation of these sections was performed blinded to the results of the tissue microarray analysis and the results of the other methodology.

Results

Data for PTEN FISH and immunohistochemistry in a subset of the CFRPTMR cohort were separately reported previously (17, 26). Briefly, of the 1275 patients with tissue sampled for the tissue microarrays, 86% (1095/1275) had evaluable PTEN status by immunohistochemistry and 14% (180/1275) had missing data (Supplementary Table S1). Of these, 17% (30/180) were missing due to ambiguous immunostaining results and 83% (150/180) had absence of tumor tissue present on the tissue microarray slides. Of the tumors with evaluable staining, 24% (258/1095) showed any PTEN protein loss, with 14% (150/1095) showing heterogeneous PTEN loss (in some but not all sampled tumor glands, best exemplified by case #10 in Figure 4A), and 10% (108/1095) showing homogeneous PTEN loss (in all sampled tumor glands). The remaining 76% (837/1095) of cases had intact PTEN protein by immunohistochemistry in all sampled tumor glands. *PTEN*FISH results were evaluable in 64% of the cases sampled on the tissue microarray (810/1275). Of the evaluable cases, *PTEN*FISH showed any *PTEN* deletion in 18% of cases, with 9% (70/810) of cases showing hemizygous deletion and 9% (75/810) of cases showing homozygous *PTEN* deletion. The remaining 82% (665/810) of cases showed two intact *PTEN* alleles.

PTEN immunohistochemistry results were available on 90% of cases with interpretable *PTEN*FISH results (731/810). The rates of *PTEN* gene and PTEN protein loss were quite similar in the subset with both FISH and immunohistochemistry results compared to the entire evaluable cohort for each assay reported separately. Overall, 22% (158/731) of cases with interpretable immunohistochemistry and FISH results showed PTEN protein loss, with 13% (96/731) showing heterogeneous loss and 8% (62/731) showing homogeneous loss. Similarly, 17% (129/731) of cases with interpretable immunohistochemistry and FISH results showed *PTEN* gene deletion (8% hemizygous and 9% homozygous).

Overall, intact PTEN immunohistochemistry was 91% specific for lack of underlying *PTEN* gene deletion. Of cases with 2 copies of the *PTEN* gene by FISH analysis, 549/602 showed intact PTEN protein (Figure 1, Tables 1 and 2). Notably, 85% (45/53) of the discrepant cases (loss of PTEN protein expression by immunohistochemistry and 2 copies of *PTEN* gene by FISH analysis) showed heterogeneous PTEN protein loss in some, but not all, sampled tumor glands, suggesting the possibility that a small area with *PTEN* deletion may have been missed in the initial FISH analysis (see below). PTEN immunohistochemistry loss was 65% sensitive for the detection of underlying hemizygous *PTEN* gene deletion since 40/62 of

cases with hemizygous *PTEN* gene deletion by FISH showed PTEN protein loss by immunohistochemistry (Figure 2). Of these cases, 65% (26/40) showed heterogeneous PTEN loss in some but not all sampled tumor glands. PTEN immunohistochemistry loss was 97% sensitive for homozygous *PTEN* gene deletion. Of cases with homozygous gene deletion by FISH, 65/67 showed PTEN protein loss by immunohistochemistry (Figure 3). Only 37% (25/67) of the cases with homozygous *PTEN* gene deletion and PTEN protein loss had heterogeneous loss of PTEN protein by immunohistochemistry. The fraction of tumors with underlying homozygous *PTEN* gene deletion differed by the extent of PTEN protein loss observed: 26% (25/96) tumors with heterogeneous PTEN protein loss had an underlying homozygous *PTEN* deletion compared to 64% (40/62) of tumors with homogeneous PTEN protein loss (p<0.0001 by Fisher's exact test).

The negative predictive value for intact PTEN immunohistochemistry was 96% (549/573) for lack of any gene deletion and 99.6% (571/573) for lack of homozygous *PTEN* deletion (Table 2). The positive predictive value of PTEN immunohistochemistry loss for presence of any *PTEN* gene deletion (homozygous or hemizygous) was 66% (105/158) overall, or 53% (51/96) for heterogeneous PTEN protein loss and 87% (54/62) for homogeneous PTEN protein loss (Table 2).

Next, we re-examined cases where there was a discrepancy between the PTEN immunohistochemistry and FISH. Overall, 53 cases with PTEN protein loss had two intact copies of *PTEN* by FISH, of which 85% (45/53) showed heterogeneous PTEN protein loss. Since only 50-100 tumor cells from two of the three tumor cores from each cases were initially evaluated by FISH, it is possible that focal tumor areas with *PTEN* gene deletion by FISH were missed or not analyzed in this blinded analysis. To examine this and other possible explanations for the immunohistochemistry-FISH discrepancy, each of these 53 discordant cases were re-reviewed for immunohistochemistry and FISH staining. Immunohistochemistry-guided FISH re-analysis in these cases revealed borderline immunohistochemistry loss in 6% (3/53) cases (Figure 4A, Case #10) and failure to analyze the identical tumor core or area by both immunohistochemistry and FISH in 34% (18/53) cases. Of the remaining discrepant cases where the immunohistochemistry result was convincing and the identical tumor area was analyzed by both methods, 41% (13/32) revealed hemizygous (n=8, Figure 4A, Case #11) or homozygous (n=5, Figure 4A, Case #12) deletion that was focal in 94% (11/13) cases and thus likely missed on initial FISH analysis. The remaining 59% (19/32) of these cases showed two copies of PTEN, thus representing truly discordant cases. One explanation for these cases is presence of a small deletion and/or mutation undetectable by FISH at one or both PTEN alleles. Another possibility is that even though the same core was evaluated by both methods in these cases, there may be heterogeneity within the core such that different levels of the core sampled on the FISH and immunohistochemistry slide may have been truly heterogeneous (Figure 4B, Case # 13). Of the two discrepant cases with homozygous PTEN deletion and intact PTEN protein, different tumor areas were analyzed in one case. In the other case, a minute focus of tumor with PTEN loss by immunohistochemistry that was initially missed was observed on re-examination (Figure 4B, Case # 14).

Finally, to further assess the effects of tumor heterogeneity on PTEN immunohistochemistry and FISH results and to determine whether this might account for discordance in some cases, we blindly studied a subset of 20 cases from the tissue microarray using standard tissue sections and compared results of immunohistochemistry and FISH on standard sections to one another and to those obtained for the tissue microarray cores of the same cases (Table 3). Cases chosen for this analysis were relatively enriched for discordance between tissue microarray-based immunohistochemistry and FISH results. In cases where the immunohistochemistry and FISH were concordant on the tissue microarray cores, results were generally highly concordant using standard sections as well. For example, in 3 cases where there was heterogeneous PTEN loss by immunohistochemistry and homozygous PTEN10ss by FISH in the tissue microarray cores, two of these tumors had clonal homozygous PTEN deletions, and the third tumor had a region with homozygous loss surrounded by a larger area with *PTEN* hemizygous loss. Similarly, in 4 cases that were PTEN intact by both immunohistochemistry and FISH on tissue microarray cores, 3 showed PTEN intact by FISH on standard sections (the fourth case failed to hybridize) and 3 showed PTEN intact by immunohistochemistry on standard sections (the fourth case showed focal PTEN loss). In some cases where there was discordance between the immunohistochemistry and FISH results on tissue microarray cores, more detailed analysis of standard sections suggested that tumor heterogeneity may be the underlying cause. Of 8 cases with heterogeneous PTEN loss by immunohistochemistry and intact PTEN by FISH on tissue microarray, 4/8 showed either hemizygous or homozygous PTEN loss by FISH on standard sections. Another case with homogeneous PTEN loss by immunohistochemistry and intact PTEN by FISH on tissue microarray revealed hemizygous PTEN loss by FISH on analysis of standard sections. Overall, these results support the possibility that underlying tumor heterogeneity is one potential cause of PTEN immunohistochemistry-FISH discordance. Despite this, tissue microarray-based evaluation of tumor PTEN status appears to be highly concordant with standard section analysis in most cases.

Discussion

There is an increasing need for validated prognostic and predictive biomarkers in prostate cancer at both ends of the clinical spectrum. Developing prognostic biomarkers to help select patients who are appropriate for active surveillance, as well as predictive biomarkers to guide the application of targeted therapy in metastatic disease remain major areas of unmet clinical need. PTEN has long been a promising marker in both regards, however, until relatively recently the lack of well validated antibodies to detect PTEN loss has made it challenging to incorporate into routine pathologic risk assessment protocols or clinical trials of PI3K-targeted agents in prostate cancer. Due to this difficulty, FISH has historically been used to assess whether PTEN is an effective prognostic biomarker by testing the association of *PTEN* gene deletion with prostate cancer progression. The results from these studies have consistently shown that *PTEN* gene deletion is associated with increased Gleason grade and stage in prostate cancer (6, 8, 10, 17, 30, 31). In addition, *PTEN* gene deletion is associated with prostate cancer progression and death in multivariable models (6-16). Though many of these previous studies have used 2-color FISH, there is increasing evidence that 4-color probes are better suited to distinguish true gene deletions from sectioning artifacts in

interphase FISH studies (Yoshimoto et al in preparation). Accordingly, our group recently demonstrated that homozygous PTEN deletion by 4-color FISH is associated with decreased recurrence-free survival in a subset of the prostate tumor cohort examined in the current study (17).

Despite these compelling data, PTENFISH has not been widely implemented in clinical prostate cancer risk stratification protocols to date for a number of reasons. First, FISH to detect gene deletions is technically challenging, requiring careful probe design (29) and rigorous cutoffs to ensure that sectioning artifacts do not result in false calls of deletion. Detecting of hemizygous deletions can be particularly challenging when nuclei are overlapping or have been distorted during preparation. Depending on tissue quality and fixation, there may also be difficulties with optimizing protease digestion such that as many as 30-40% of cases cannot be evaluated on the first attempt when using tissue microarrays, though this may be less of an issue for biopsies (17). In large part because it is so technically challenging, FISH is relatively expensive compared to immunohistochemistry, and it has been harder to integrate the daily workflow of pathology laboratories as a reflexive test. Finally, though *PTEN* is most commonly lost via larger genomic deletions in prostate cancer, as many as 10-20% of cases may have mutations, small insertions or deletions that are not detectable by FISH, in addition to potential epigenetic and miRNA-mediated mechanisms of PTEN loss (1-5, 32). To address these challenges, several groups have developed immunohistochemistry assays to query PTEN status in tissue (9, 19, 33). While a number of such assays have been published, for the most part, these assays have largely been compared to 2-color FISH in only small scale studies with around 100 tumors each (23, 24, 34, 35). In the only large studies to compare immunohistochemistry and FISH, there was only weak (κ =0.5) (14) or no significant correlation (13) between the assays, suggesting a failure of the immunohistochemistry and/or FISH assay to analytically validate.

We used a commercially available rabbit monoclonal antibody to develop an immunohistochemistry assay to assess PTEN protein loss in prostate cancer and showed that this assay is reasonably sensitive for detection of *PTEN* gene deletion by 2-color FISH or high density SNP array in prostate cancer samples and shows minimal inter-observer variability in interpretation (9). Similarly, the assay performed well versus 4-color FISH in a small cohort of needle biopsy specimens (28). Using this assay, our group previously demonstrated that PTEN protein loss is associated with an increased risk of recurrence and progression in surgically treated cohorts of prostate cancer patients (11, 12).

To facilitate clinical use of the assay, we adapted it to the automated Ventana staining platform with clinical-grade reagents suitable for *in vitro* diagnostic use. This assay was clinically validated in a recent study showing that PTEN loss is associated with increased risk of lethal prostate cancer in a large population-based cohort in multivariable models (18). Despite a 4-category scoring system, the assay has shown high inter-observer reproducibility in a number of cohorts (including the current one), with κ values exceeding 0.9 (18, 26). In the current study, we analytically validated this automated assay by comparing it to 4-color *PTEN*FISH across a large multi-institutional cohort of prostate cancer patients. Remarkably, we found that the automated immunohistochemistry assay was 91% specific for 2 intact copies of the *PTEN* gene and 97% sensitive for homozygous *PTEN* gene deletions. This is

by far the highest sensitivity and specificity reported for a PTEN immunohistochemistry assay relative to FISH. This improved sensitivity and specificity is in part due to the improved specificity of the automated immunohistochemistry assay versus the manual assay and also due to the improved 4-color FISH assay which uses two *PTEN* gene flanking probes, in addition to a centromeric control and a *PTEN* probe to detect *PTEN* gene deletions. Surprisingly, the immunohistochemistry assay was also 65% sensitive for detection of hemizygous *PTEN* gene deletion, suggesting that there is complete protein loss in a large fraction, perhaps even a majority, of apparently hemizygous cases. This is most likely due to truncating mutations (nonsense, frameshift and splice site mutations) or epigenetic modifications at the second allele that are undetectable by FISH yet lead to protein loss (1, 3, 5, 36). Interestingly, though the prevalence of such mutations in *PTEN* is below 5% in most prostate tumors, many of these mutations are truncating alterations occurring in cases with hemizygous deletions that would lead to protein loss detectable by immunohistochemistry (1-5).

In addition to the potential increased sensitivity of immunohistochemistry versus FISH for detecting combinations of events including copy loss, point mutations, small insertions and deletions and epigenetic modifications leading to *PTEN* inactivation, immunohistochemistry is also very useful for screening for areas of focal PTEN loss. By necessity, *PTEN*FISH is analyzed at high magnification, examining 50-100 nuclei, which may miss small areas of loss within the sampled tumor. In contrast, immunohistochemistry can be easily screened at low magnification and still afford a nearly cell-by-cell resolution image of PTEN expression. In the current study, in over 40% of cases where PTEN immunohistochemistry detected loss and *PTEN*FISH was initially read as 2 copies in the identical tumor core, rescreening the FISH guided by areas of immunohistochemistry loss resulted in detection of small areas with *PTEN* deletion, initially missed or beneath the cutoff for the FISH scoring. This result, in addition to the high negative predictive value of intact immunohistochemistry for lack of deletion strongly suggests that immunohistochemistry screening for PTEN loss is likely to be an efficient and cost-effective strategy to ascertain PTEN status in tissue sections.

Akin to HER2 assessment in breast, it is ultimately likely that the best protocol will be to perform reflexive FISH on a subset of prostate tumors after initial immunohistochemistry screening. Clearly, in cases with ambiguous immunohistochemistry results (<5%), FISH will have an important role. However, there may also be a role for FISH in cases with heterogeneous loss of PTEN by immunohistochemistry. As in previous cohorts (12), in the current cohort we found that homogeneous PTEN immunohistochemistry loss was more strongly associated with decreased recurrence-free survival compared to heterogeneous PTEN protein loss in both univariate and multivariate analyses (26). The explanation for why focal PTEN loss is a less potent prognostic indicator than homogeneous loss remains unclear. Homogeneous PTEN loss may be a surrogate indicator for expansion of a single, dominant clone of tumor cells. Alternatively, perhaps loss of PTEN in a larger number of cells increases risk of tumor progression for stochastic reasons. Finally, this result may also be related to the higher prevalence of homozygous *PTEN* deletion among the cases with homogeneous immunohistochemistry loss, compared to the cases with heterogeneous immunohistochemistry loss (64% vs 26%; p<0.0001 by Fisher's exact test). Indeed, in the subset of the current cohort where *PTEN*FISH was correlated with disease outcomes, only

homozygous but not hemizygous *PTEN* loss was associated with decreased recurrence free survival in multivariate models (17). Thus, it may be that tumors with heterogeneous PTEN protein loss and underlying homozygous *PTEN* gene deletion have outcomes roughly equivalent to cases with homogeneous PTEN protein loss (the majority of which have homozygous deletion). Though larger case numbers than were included in the current study will be required to formally address this hypothesis, this would suggest that it may useful to perform reflexive FISH in the case of heterogeneous PTEN protein loss by immunohistochemistry (14% of total cases in current cohort) to determine whether there is underlying homozygous *PTEN* gene deletion. The FISH could be guided by the immunohistochemistry to focus on areas with protein loss, increasing the sensitivity of the assay in this way.

There are a number of limitations of the current study. Though both FISH and immunohistochemistry were performed on the same tissue microarrays, analysis of all tissue microarray cores was not technically feasible for both methods in all cases and correlation between the two assays was done on a tumor-by-tumor rather than core-by-core basis for most cases. Thus, some of the disagreements between FISH and immunohistochemistry likely came about because of tumor heterogeneity, where different areas of the same tumor were being analyzed by each assay, and standard section analysis of a subset of cases largely bears this out. In addition, the gold-standard for assessing PTEN gene status is not clear at this point. Though FISH can detect larger deletions which are the most common mechanism of loss in prostate cancer, it will miss smaller deletions, as well as indels and missense mutations which may inactivate the gene. Thus, in cases where the same tumor tissue was analyzed, it is impossible to know the true cause of the apparent discrepancies between FISH and immunohistochemistry without using a third methodology such as sequencing to examine for gene alterations that would be missed by FISH (these studies are ongoing in separate cohorts currently). Finally, due to the relatively small numbers of discordant cases overall, it was not feasible to do a meaningful analysis comparing FISH and immunohistochemistry for prediction of prognosis in these cases, to determine which assay is a better prognostic tool.

In conclusion, in a large multi-institutional cohort of prostate tumors, our immunohistochemistry assay for PTEN loss shows the highest specificity and sensitivity for *PTEN* gene deletion reported for an immunohistochemistry assay to date. These data strongly suggest that immunohistochemistry is a cost-efficient method to screen for PTEN loss in prostate tumors, requiring ~\$100 and a single 4 µm tumor section for assay performance. In cases with ambiguous PTEN immunohistochemistry results or heterogeneous PTEN protein loss, reflexive *PTEN*FISH may be a useful confirmatory test. This inexpensive, automated and analytically validated immunohistochemistry assay has already been used to demonstrate the association of PTEN loss with lethal prostate cancer in a large population-based cohort in multivariable models (18). Ultimately, its portability will enable the performance of clinical validation studies on a large number of additional cohorts, credentialing PTEN as a prognostic and potentially predictive biomarker in diverse clinical settings.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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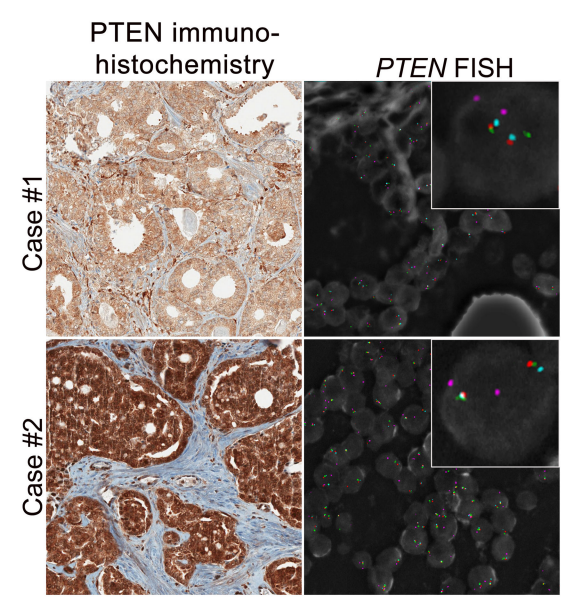


Figure 1.

Prostate cancer cases showing intact PTEN protein with 2 intact PTEN gene alleles. <u>Cases</u> <u>#1 and 2</u>: PTEN immunohistochemistry demonstrates intact PTEN protein (left), while fourcolor FISH image from adjacent section (right) shows two intact *PTEN* alleles (see enlarged inset—two red signals) with two intact copies flanking genes, *WAPAL* (green) and *FAS* (aqua) as well as chromosome 10 centromeres (pink). Author Manuscript

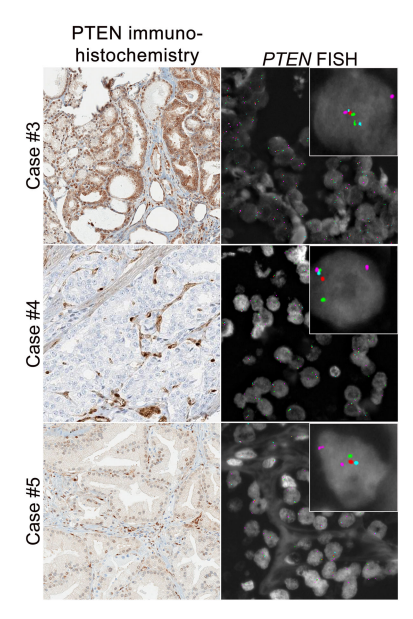


Figure 2.

Prostate cancer cases showing variable PTEN protein expression with hemizygous *PTEN* gene deletion. <u>Case #3</u>: PTEN immunohistochemistry demonstrates intact PTEN protein (left), with four-color FISH image from an adjacent section showing a hemizygous *PTEN* deletion with loss of one *PTEN* gene (see enlarged inset-one red signal). Since both centromeres (pink) and the *WAPAL* (green) and *FAS* (aqua) probes that flank either side of *PTEN* are retained it is likely that this hemizygous deletion is interstitial and restricted to the *PTEN* region. <u>Case #4</u>: PTEN immunohistochemistry image shows homogeneous loss of PTEN protein (left) while FISH image from an adjacent section (right) shows a hemizygous *PTEN* deletion (see enlarged inset-one red signal). Concurrent hemizygous deletion of the adjacent *FAS* gene probe (one aqua signal missing) but retention of two copies of the centromere and *WAPAL* gene probes indicates the deletion includes both the *PTEN* and *FAS* genes. <u>Case #5</u>: PTEN immunohistochemistry image shows somewhat light, but intact

immunostaining for PTEN protein (left) while the FISH image from an adjacent section (right) shows a hemizygous *PTEN* deletion (see enlarged inset-one red signal). Since there was concurrent loss of the *WAPAL*, *PTEN* and *FAS* gene probes (green, red and aqua, respectively), but retention of both centromeres (pink), this hemizygous deletion extends outside the *PTEN* region in both directions.

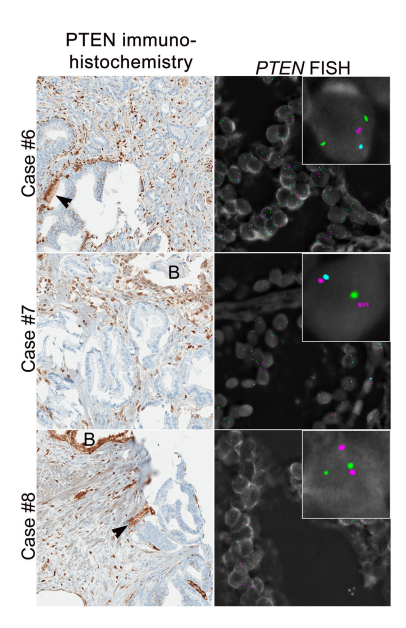


Figure 3.

Prostate cancer cases showing absence of PTEN protein expression with homozygous *PTEN* gene deletion. <u>Case #6</u>: PTEN immunohistochemistry image (left) shows loss of PTEN protein in tumor glands. Intraductal spread of tumor is present in this case and retention of PTEN protein is seen in benign basal and luminal cells of duct containing tumor (arrowhead). Four-color FISH image from an adjacent section (right) shows a homozygous deletion with loss of both *PTEN* genes (see enlarged inset - no red signals). The retention of the centromeres (pink) and both *WAPAL* genes (green), but the presence of only one copy of the *FAS* gene (aqua) indicates that one of the deletions involved both the *PTEN* and *FAS* genes. <u>Case #7</u>: PTEN immunohistochemistry image (left) shows loss of PTEN protein in tumor glands, with retention in entrapped benign gland (B). FISH image from an adjacent section (right) shows a homozygous *PTEN* deletion (see enlarged inset - no red signals). The retention of the centromeres (pink) but concurrent loss of one *WAPAL* (green) and one *FAS*

gene (blue) indicates the deletions extend outside the *PTEN* region. <u>Case #8</u>: PTEN immunohistochemistry image (left) shows loss of PTEN protein in tumor glands, with retention in adjacent benign gland (B) and nearby endothelial cells (arrowhead). (FISH image from an adjacent section (right) shows a homozygous *PTEN* deletion (see enlarged inset - no red signals). The retention of the centromeres and both the *WAPAL* genes (green), but the concurrent loss of both *FAS* (blue) and *PTEN* (red) indicates that both copies of chromosomes 10 have deletions involving these genes.

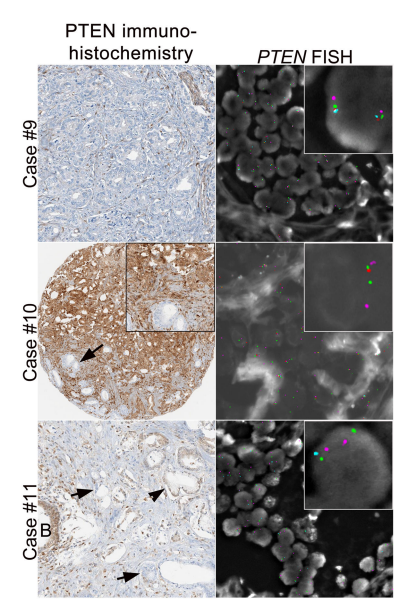


Figure 4A.

Prostate cancer cases with discordant PTEN immunohistochemistry and FISH results on initial review. <u>Case #9</u>: PTEN immunohistochemistry demonstrates very weak cytoplasmic immunostaining with loss of nuclear immunostaining and thus was called negative on initial review, though in retrospect it may be better classified as ambiguous due to weak staining and absence of benign glands for comparison (left). Four-color FISH image from an adjacent section that is representative of all examined cores in this tissue microarray (right) indicates that the *PTEN* gene does not have a detectable deletion by FISH. The enlarged inset shows that the centromeres, *WAPAL*, *PTEN* and *FAS* gene probes are each present as two copies. <u>Case #10</u>: PTEN immunohistochemistry image (left) shows heterogeneous PTEN loss in some tumor glands (arrow) but PTEN protein is expressed by majority of other tumor glands in this core. FISH image from an adjacent section (right) was initially read as PTEN intact, but shows a focal area with hemizygous *PTEN* deletion recognized on re-examination

guided by immunohistochemistry. The enlarged inset shows there is only one copy of the red *PTEN* gene probe (one red signal) and loss of both aqua *FAS* gene probes. <u>Case #11</u>: PTEN immunohistochemistry image (left) demonstrates heterogeneous PTEN loss in some tumors glands (arrows) but not in others (arrowheads). FISH image from an adjacent section (right) shows the small area of the section that had a homozygous *PTEN* deletion on re-examination. The enlarged inset shows that there are no copies of the red *PTEN* gene probe and one copy of the aqua *FAS* gene probe, but retention of the adjacent *WAPAL* and centromere probes.

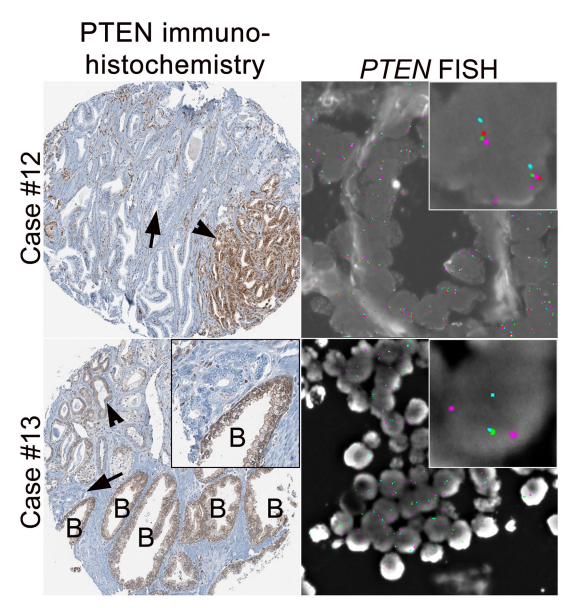


Figure 4B.

Prostate cancer cases with discordant PTEN immunohistochemistry and FISH results on initial review. <u>Case #12</u>: PTEN immunohistochemistry image (left) shows heterogeneous loss of PTEN protein in some tumor glands (arrow) but not in others (arrowhead). A FISH image from an adjacent section that is representative of all examined cores in this tissue microarray (right) indicates that the *PTEN* gene does not have a detectable deletion by FISH. The enlarged inset shows that the centromeres, *WAPAL, PTEN* and *FAS* gene probes are each present as two copies. The heterogeneous loss in this case may have resulted in different tumor areas sampled in slides for immunohistochemistry and that for FISH. <u>Case #13</u>: PTEN immunohistochemistry image (left) shows predominantly intact/light immunostaining in tumor glands (arrowhead) and benign glands (B) with a very focal area of tumor with PTEN loss identified on re-review after FISH analysis (arrowhead, inset).

PTEN deletion. The enlarged inset shows that there are no copies of the red *PTEN* gene probe and loss of one green *WAPAL* gene probe but retention of both the *FAS* and the centromere probes.

Table 1

Summary of PTEN immunohistochemistry by PTEN FISH status.

| | | | PT | PTEN FISH | | |
|---------------------------|--------|--------|--------|---------------|---------------|----------|
| | Intact | act | Hemi-d | Hemi-deletion | Homo-deletion | leletion |
| | Z | % | N | ⁰% | Ν | ⁰‰ |
| PTEN immunohistochemistry | | | | | | |
| Intact | 549 | 549 91 | 22 | 35 | 2 | 3 |
| Heterogeneous loss | 45 | L | 26 | 42 | 25 | 37 |
| Homogeneous loss | 8 | 1 | 14 | 23 | 40 | 60 |

Table 2

Performance metrics for PTEN immunohistochemistry compared to gold-standard *PTEN* FISH

| | | n |
|-------------------------------------|----|---------|
| Specificity | 91 | 549/602 |
| Sensitivity for homozygous deletion | 97 | 65/57 |
| Sensitivity for hemizygous deletion | 65 | 40/62 |
| Positive predictive value | 66 | 105/158 |
| Negative predictive value | 96 | 549/573 |

Table 3

Comparison of PTEN immunohistochemistry and FISH results on tissue microarray cores and standard tissue section slides

| Case | Tissue microarray PTEN immunohistochemistry | standard slide PTEN immunohistochemistry | Tissue microarray PTEN FISH | standard slide PTEN FISH |
|------|--|---|--------------------------------|----------------------------------|
| 1 | intact | intact | intact | intact |
| 2 | intact | heterogeneous loss | intact | intact |
| 3 | intact | intact | intact | intact |
| 4 | intact | intact | intact | failure |
| 5 | heterogeneous loss | heterogeneous loss | homo-deletion | hemi-deletion and homo-delletion |
| 6 | heterogeneous loss | heterogeneous loss | homo-deletion | homo-deletion |
| 7 | heterogeneous loss | heterogeneous loss | homo-deletion | homo-deletion |
| 8 | heterogeneous loss | heterogeneous loss | intact | intact |
| 9 | heterogeneous loss | heterogeneous loss | intact | intact |
| 10 | heterogeneous loss | intact | intact | intact |
| 11 | heterogeneous loss | heterogeneous loss | intact | intact |
| 12 | heterogeneous loss | intact | intact | hemi-deletion |
| 13 | heterogeneous loss | heterogeneous loss | intact | homo |
| 14 | heterogeneous loss | heterogeneous loss | intact | homo |
| 15 | heterogeneous loss | heterogeneous loss | intact | hemi-deletion of WAPAL |
| 16 | homogeneous loss | homogeneous loss | hemi-deletion | hemi-deletion |
| 17 | homogeneous loss | heterogeneous loss | intact | hemi-deletion |
| 18 | ambiguous | intact | intact | intact |
| 19 | heterogeneous loss | heterogeneous loss | core missing | intact |
| 20 | heterogeneous loss | heterogeneous loss | core missing | intact |