Purpose: To assess the underlying genomic variation of prostate gland microenvironments of patients with prostate adenocarcinoma in the context of colocalized multiparametric magnetic resonance (MR) imaging and histopathologic assessment of normal and abnormal regions by using whole-exome sequencing.

Materials and Methods: Six patients with prostate adenocarcinoma who underwent robotic prostatectomy with whole-mount preservation of the prostate were identified, which enabled spatial mapping between preoperative multiparametric MR imaging and the gland. Four regions of interest were identified within each gland, including regions found to be normal and abnormal via histopathologic analysis. Whole-exome DNA sequencing (>50 times coverage) was performed on each of these spatially targeted regions. Radiogenomic analysis of imaging and mutation data were performed with hierarchical clustering, phylogenetic analysis, and principal component analysis.

Results: Radiogenomic multiparametric MR imaging and whole-exome spatial characterization in six patients with prostate adenocarcinoma (three patients, Gleason score of 3 + 4; and three patients, Gleason score of 4 + 5) was performed across 23 spatially distinct regions. Hierarchical clustering separated histopathologic analysis–proven high-grade lesions from the normal regions, and this reflected concordance between multiparametric MR imaging and resultant histopathologic analysis in all patients. Seventy-seven mutations involving 29 cancer-associated genes across the 23 spatially distinct prostate samples were identified. There was no significant difference in mutation load in cancer-associated genes between regions that were proven to be normal via histopathologic analysis (34 mutations per sample ± 19), mildly suspicious via multiparametric MR imaging (37 mutations per sample ± 21), intermediately suspicious via multiparametric MR imaging (31 mutations per sample ± 15), and high-grade cancer (33 mutations per sample ± 18) (P = .30). Principal component analysis resolved samples from different patients and further classified samples (regardless of histopathologic status) from prostate glands with Gleason score 3 + 4 versus 4 + 5 samples.

Conclusion: Multiregion spatial multiparametric MR imaging and whole-exome radiogenomic analysis of prostate glands with adenocarcinoma shows a continuum of mutations across regions that were found via histologic analysis to be high grade and normal.

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Online supplemental material is available for this article.
Prostate cancer is the most prevalent malignancy in men and third most common cause of death because of malignancy (1). The phenotype and course of the disease span a broad spectrum, from an indolent tumor to an aggressive metastatic carcinoma (2,3). Whereas there continue to be advancements in screening and treatment, some of the basic underlying biologic processes regarding gland changes over decades of life have yet to be elucidated.

The recognition and appreciation of neoplasia as an evolutionary process has long been recognized (4–6). As technological capabilities have advanced there has been deeper characterization of tumors by histopathologic analysis, clinical imaging, and high-throughput data analysis that has shed further light on mechanisms of development, progression, and differences in tumor grade and aggressiveness (7,8). Intratumoral heterogeneity has been highlighted as a feature of multiple malignancies (9); however, variation in the imaging appearance, histopathologic analysis, and whole-exome variation within tumors and their surrounding microenvironments proven to be normal via histopathologic analysis have not, to our knowledge, been explored.

In recent years, radiogenomics has become an increasingly popular approach for linking molecular phenotyping with clinical imaging (10–17). Simultaneous characterization of microscopic multiparametric magnetic resonance (MR) imaging features with microscopic genomic signatures in the context of well-defined histopathologic structure in a spatially colocalized manner may further inform the underlying variation of microenvironments in prostate glands that developed adenocarcinomatous nodules.

We set forth to further evaluate these features simultaneously in prostate glands with confirmed adenocarcinoma by histopathologic analysis (Gleason score, >6). We leveraged the colocalization ability of preserved whole-mount prostates, resected for treatment of adenocarcinoma, with preoperative multiparametric MR imaging to selectively interrogate multiple regions of interest, including regions determined via histopathologic analysis to be abnormal as well as normal by using whole-exome DNA sequencing (sequencing all of the expressed genes in the human genome). The purpose of this study was to assess the underlying genomic variation of prostate gland microenvironments in patients with prostate adenocarcinoma in the context of colocalized multiparametric MR imaging and histopathologic assessment of normal and abnormal regions by using whole-exome sequencing.

### Materials and Methods

#### Patients and Materials

This was a Health Insurance Portability and Accountability Act–compliant, institutional review board–approved study that involved tissue and multiparametric prostate MR examinations in six retrospectively identified patients with a diagnosis of prostate adenocarcinoma and Gleason scores greater than or equal to 7 (3+4) between July 2009 and October 2011. Selection criteria was on the basis of acquisition of preoperative diagnostic multiparametric MR examinations in addition to paraffin-embedded whole mounts of the resected prostates with adequate tissue for DNA extraction from areas determined via histopathologic analysis to be normal and abnormal, and normal but with appearance at MR imaging that was mildly or moderately suspicious for cancer. Initially, 104 patients were identified with preoperative multiparametric MR imaging who were then cross-matched with available tissue whole mounts at the Translational Pathology Core Laboratory at University of California, Los Angeles, with confirmed adenocarcinoma and at least one other region at MR imaging noted to be abnormal, reducing the list to 14 patients of which one was Gleason score 3+3. The remaining 13 patients were evaluated...
on the basis of availability of adequate quantities of tissue for whole-exome sequencing, which resulted in the six glands ultimately analyzed in this study.

Preoperative 3-T pelvic multiparametric MR imaging (Siemens, Erlangen, Germany) with endorectal coils was performed on all patients. The examinations included T1-weighted imaging, T2-weighted turbo spin-echo imaging, diffusion-weighted imaging, T2 water and fat-saturated three-dimensional spectroscopic imaging, and intravenous contrast material–enhanced T1-weighted gradient-echo fat-saturated images before and after administration of gadolinium chelate (Table 1). Imaging examinations were reviewed by an abdominal and genitourinary radiologist (D.J.M., with 10 years of experience). Objective and subjective assessment of the glands was performed through standardized assessment of T2 signal, estimation of the average apparent diffusion coefficients, pharmacokinetic enhancement curve characteristics from dynamic contrast–enhanced images, and spectroscopic imaging (18), with prospective evaluation on a five-point ranked categorical scale as previously described by Le et al (19) and is the standard of care at our institution.

Each prostate gland was sectioned at 4–5 mm intervals. By using a whole-mount technique, tissue slices were then formalin fixed, paraffin embedded, and microtome cut (20,21). Gleason scoring was performed according to the 2005 International Society of Urological Pathology consensus recommendations (22).

Part of the motivation of this study was to characterize focal areas of the prostate, or microenvironments, that showed concordance between histopathologic analysis and multiparametric MR imaging in addition to characterizing regions that were normal by histopathologic appearance but demonstrated some imaging abnormalities that were not suspicious for cancer by Prostate Imaging Reporting and Data System (PI-RADS)–like criteria (23,24). For example, a focal nodule in the peripheral zone may exhibit decreased T2 signal intensity, an apparent diffusion coefficient of 1300 mm²/sec, but no abnormal enhancement and normal spectroscopy; this would not warrant tissue sample by clinical standards, but would be included in the “mildly suspicious” category. Four different regions of interest were targeted within each gland from the whole-mount paraffin-embedded surgically resected prostates on the basis of the combination of pathologic and imaging criteria to construct the following imaging score matrix (Appendix E1 [online]):

<table>
<thead>
<tr>
<th>Pulse Sequence</th>
<th>Orientation</th>
<th>TR/TE (msec)</th>
<th>FOV (mm)</th>
<th>Matrix</th>
<th>Thickness (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3D TSE T2 (SPACE)</td>
<td>Axial</td>
<td>3800/101</td>
<td>140</td>
<td>256 × 256</td>
<td>1.5</td>
</tr>
<tr>
<td>2D TSE T2</td>
<td>Axial, Coronal</td>
<td>3800/101</td>
<td>200</td>
<td>320 × 320</td>
<td>3.6</td>
</tr>
<tr>
<td>Echo planar DWI</td>
<td>Axial</td>
<td>3900/60</td>
<td>210 × 260</td>
<td>130 × 160</td>
<td>3.6</td>
</tr>
<tr>
<td>Dynamic contrast enhancement (TWIST)</td>
<td>Axial</td>
<td>3.9/1.4</td>
<td>260</td>
<td>160 × 160</td>
<td>3.6</td>
</tr>
</tbody>
</table>

Note.—The parameters were as follows: three-dimensional turbo spin-echo T2-weighted imaging (SPACE), echo train length of 13; two-dimensional turbo spin-echo T2-weighted imaging, echo train length of 13–25; echo-planar diffusion-weighted imaging, b values of 0, 100, 400, and 800 sec/mm³; dynamic contrast–enhanced (time-resolved angiography with interleaved stochastic trajectories) imaging, 12° flip angle, 4.75 seconds per acquisition over 5 minutes; MR signal intensity, 1300 Hz, six averages, 256 spectral points. 2D = two dimensional, 3D = three dimensional, DWI = diffusion-weighted imaging, FOV = field of view, SPACE = sampling perfection with application optimized contrasts using different flip-angle evolution, TE = echo time, TR = repetition time, TSE = turbo spin echo, TWIST = time-resolved angiography with interleaved stochastic trajectories.

Thus, each gland was re-evaluated by a genitourinary pathologist (J.I.L., with >20 years of experience in genitourinary pathologic analysis) to confirm the initial diagnosis and to also identify appropriate regions to obtain tissue samples from (ie, one region within the high-grade area and three areas that were normal-appearing by histopathologic analysis but still within the peripheral zone of the gland).

Matching was performed during scheduled dedicated sessions where the pathologist, radiologist, and research staff would review the MR image and whole-mount histopathologic analysis to determine whether a finding at MR imaging constituted a match on the whole mount, as part of clinical quality assurance, with mismatches characterized as false-negative findings (ie, cancers that were not prospectively identified at MR imaging) and false-positive findings (ie, lesions on MR images that did not correspond to cancer). Therefore, the prospectively identified regions of interest on MR images with their corresponding categorical and quantitative assessments were correlated with cancerous (or benign) regions on the whole mount.

Isolated tissue samples were provided to Otogenetics (Otogenetics Corporation, Atlanta, Ga) for whole-exome DNA sequencing with Illumina HISEquation 2000 sequencers (Illumina, San Diego, Calif) with paired-end reads of 90–100 nucleotides, greater than or equal to 50 times coverage. The “moderately suspicious” tissue sample from patient 3 was of insufficient quality to sequence, resulting in 23 samples with whole-exome sequencing in total from six patients. Intron and nonsynonymous mutations were excluded. Furthermore,
we only included mutations that were detected in at least two of the 23 independent samples. To identify mutations that were more likely to be related to the underlying malignancy within the tumors, we focused our analysis by using a list of curated genes in the literature that have been associated with cancer (25). The gene set of interest was reduced by finding the intersection with the Catalogue of Somatic Mutations in Cancer database (25). The mutation profiles were used to define mutation score matrices (Appendix E1 [online]).

Statistical Analysis

We performed the Kruskal-Wallis test to assess any statistically significant variations among the tissue sample physical locations, tissue classification, Gleason scores, chromosomal location, and mutation profiles. We performed two-way hierarchical clustering with complete linkage by using the Euclidean norm on the imaging score matrix and the mutation score matrix. A phylogenetic tree was constructed from the pairwise distances between the samples on a mutation-by-mutation basis. We calculated the cophenetic coefficient for the entire tree and performed principal component analysis of the signed binary mutation matrix. We analyzed the highest ranking principal components, which accounted for at least 50% of the variation across all of the samples. Assessment of the most significant genes contributing to the third principal component was assessed by selecting the genes with absolute value greater than 0.4. The Student t test was performed to assess any statistically significant age difference between the Gleason score subgroups. Pearson correlation coefficients were calculated between the ordinal imaging scores and the mutation (weighted and binary) matrices on a sample-by-sample basis. Multiple hypothesis correction was performed by using the Bonferroni method. Data processing, calculations, and figure generation were performed by using software (Python v2.7, Matlab 2015b, MathWorks, Natick, Mass; and R v3.1.1, R Project for Statistical Computing, www.r-project.org).

Table 2

Summary of Patient Characteristics

<table>
<thead>
<tr>
<th>Patient No.</th>
<th>Age (y)</th>
<th>PSA (ng/mL)</th>
<th>Gleason Score</th>
<th>Stage</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>70</td>
<td>7.1</td>
<td>4 + 5</td>
<td>T3a, NX, MX</td>
</tr>
<tr>
<td>2</td>
<td>70</td>
<td>0.8</td>
<td>4 + 5</td>
<td>T3a, N0, MX</td>
</tr>
<tr>
<td>3</td>
<td>68</td>
<td>6.1</td>
<td>3 + 4</td>
<td>T2c, N0, MX</td>
</tr>
<tr>
<td>4</td>
<td>68</td>
<td>6.1</td>
<td>4 + 5</td>
<td>T2c, N0, MX</td>
</tr>
<tr>
<td>5</td>
<td>74</td>
<td>18.9</td>
<td>3 + 4</td>
<td>T3a, N0, MX</td>
</tr>
<tr>
<td>6</td>
<td>66</td>
<td>9.5</td>
<td>3 + 4</td>
<td>T3a, N0, MX</td>
</tr>
</tbody>
</table>

Note.—There was no statistically significant difference between the ages of the patients. Elevated prostate-specific antigen results from screening examinations prompted referral for multiparametric MR imaging for all patients except patient 2, who was found to have a palpable nodule at physical examination and was referred for imaging evaluation despite a normal prostate-specific antigen.

Please refer to the Appendix E1 (online) for additional details.

Results

Patients

Six patients with prostate adenocarcinoma were identified (three patients with Gleason score of 3+4; mean age, 68 years [age range, 66–74 years]; and three patients with Gleason score of 4+5; mean age, 68 years [age range, 68–70 years]). There was no statistically significant difference between the two groups on the basis of age (Student t test, \( P > .999 \)) (Table 2).

Multiparametric MR Imaging

The whole-mount prostate specimens in conjunction with three-dimensional anatomic MR imaging enabled tissue samples to be analyzed from focal regions, or microenvironments, of the glands by using imaging, and it enabled histopathologic analysis–based characterization (Fig 1). Regions with combinations of T2 shortening, impaired diffusion, abnormal enhancement characteristics, and/or elevated choline were concordant with high-grade regions proven with histologic analysis (Fig 1; Tables E1, E2 [online]; Appendix E1 [online]), consistent with findings in the literature (4,26,27).

Unsupervised hierarchical clustering of multiparametric MR imaging features (T2 signal, apparent diffusion coefficient, contrast enhancement curve, and spectroscopy) across all samples and patients revealed clustering of samples according to normal and histopathologic groups (Fig 2). As expected, normal multiparametric MR imaging samples were in a subcluster and nodules suspicious for cancer (ie, high) were in a separate subcluster. The mild regions clustered with the normal regions of interest and the moderate regions clustered with the nodules suspicious for cancer. The separation of regions of interest that were normal and suspicious for cancer is consistent with observations in the literature and PI-RADS–based classification for radiology-pathologic (known as rad-path) correlations between multiparametric MR imaging and histologic analysis differentiating normal tissue from tumor.

Whole-Exome DNA Sequencing

Global assessment of single nucleotide variants revealed 94166 cumulative variant calls across all samples (Fig 3). The majority of mutations were nonsynonymous mutations (97.2% [91578 of 94166]), with only 1.5% (1380 of 94166) frameshift mutations and 1.3% (1199 of 94166) stop-gain or stop-loss mutations identified. There was no evidence of bias or significant association between the different microenvironments and the chromosomal location of the mutations (Kruskal-Wallis test, \( P = .268 \)) or with the different microenvironments and type of mutation (eg, frameshift, nonsynonymous, stop gain, stop loss).

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Figure 1: Mapping between three-dimensional prostate whole mounts and multiparametric MR imaging. Tissue sections from whole-mount slides correspond to different regions of interest: adenocarcinoma (Gleason score, \(\geq 3+4\)) in red outline; moderately suspicious according to multiparametric MR imaging in blue outline; and mildly suspicious by multiparametric MR imaging in yellow outline. The nodule of interest (arrowheads) is shown in each of the fields of view, which are, from left to right: axial T2-weighted image, apparent diffusion coefficient map, spectroscopy profile, and coronal T2-weighted images (which map to the corresponding axial multiparametric MR imaging images and histologic sections), and annotated whole-mount section. The red outlined images highlight a cancer nodule in the right apical posterior peripheral zone: adenocarcinoma with radiologic and pathologic-analysis concordance with abnormal T2 signal, positive diffusion restriction with apparent diffusion coefficient of 1015 mm\(^2\)/sec, and abnormal spectroscopy (ie, reversal of choline and creatine-to-citrate ratio) that was sectioned and whole-exome sequenced according to the dotted black line in the tissue section (right-hand side). The blue outlined images depict a lesion moderately suspicious for cancer (that was normal according to histopathologic analysis) but with abnormal hypointense T2 signal, apparent diffusion coefficient of 1347 mm\(^2\)/sec reflecting some restricted diffusion, and normal spectroscopy. The yellow outlined images highlight a histologically normal nodule with abnormal T2 signal hypointensity but no evidence or diffusion or spectroscopic abnormalities.

Figure 2: One-way hierarchical clustering of multiparametric features across 23 regions of interest in six patients show radiologic and pathologic (path) concordance. Red colored blocks reflect abnormal feature values (eg, decreased T2 signal intensity, decreased diffusion, wash-in and/or wash-out enhancement characteristics, and abnormal spectroscopy) and yellow blocks reflect normal parameter measurements. The two principle branches of the dendrogram across the x-axis segregate the samples into normal and abnormal according to histopathologic analysis, with multiparametric MR imaging classified samples mildly and moderately suspicious for cancer intermixed among the two principle groups. \(K_{\text{trans}}\) = volume transfer constant.
Next, we focused the analysis on the subset of curated cancer-associated genes from the Catalogue of Somatic Mutations in Cancer database; removal of noncancer-associated mutations was accompanied by a decrease in variation between individual tissue samples, which suggested that the vast majority of variant calls, which were filtered out, were likely passenger mutations (Figs E3, E4 [online]; Table E3 [online]). There was no significant variation in cancer-associated variants (mutations) among the four different groups (histopathologic analysis–confirmed normal, 34 mutations per sample ± 19 [standard deviation]; multiparametric MR imaging–confirmed mild suspicion, 37 mutations per sample ± 21; multiparametric MR imaging–confirmed moderate to high suspicion, 34 mutations per sample ± 19; multiparametric MR imaging–confirmed high suspicion, 34 mutations per sample ± 19).

Figure 3: Overview of cumulative nonintronic, annotated mutations across all 23 samples. The majority of mutations were nonsynonymous (NS) with less than 3% and composed of frameshift, stop-gain, or stop-loss mutations. Each subpanel from left to right summarizes the 94,166 mutations, starting with the first mutation in the upper left and the last mutation in the lower right of each vertical panel. The subpanels consist of 94,166 small rectangular boxes that correspond to individual mutations, the coordinates of each subrectangle within each column correspond to the same mutation across all panels (except for the coverage panel, which summarizes the average across each corresponding row). NC = noncoding, UTR = untranslated region. Visually there was no evidence of significant biases between the different microenvironments and the location or types of mutations. There is the suggestion of significant variation in mutation coverage for patient 3, which was confirmed statistically (Kruskal-Wallis test, \( P < 2 \times 10^{-16} \)).
Figure 4: Two-dimensional hierarchical clustering of data of mutations versus the colocalized genomic-sampled regions show clustered subgroups that correspond to individual patients. Row dendrogram (left side) corresponds to clustering of the colocalized genomic-sampled regions and are colored coded by patient and similarly labeled by patient number, tissue location, and classification (ie, normal, mild, moderate, and path) on the right. Samples clustered most strongly by groups of individual patients, as color coding of the patient specific subbranches reflect. Column dendrogram (top) reveals clustering of filtered and annotated mutations. Yellow reflects the presence of a mutation and black reflects an absence. Additional detailed annotations of the mutations are provided in Tables E4 and E5 (online).

Radiogenomic Assessment

Phylogenetic analysis of the complete set of samples revealed a branching pattern that grouped together by patient at the most distal end (ie, leafs), although grouping according to groups of mutations was noted more proximally (Fig 5). The cophenetic correlation across the entire tree was 0.91. The branching pattern of the tree revealed the highest concordance among independent tissue samples from different regions of the prostate that were acquired from the same patient, which provided a degree of validation of the mutations. Tissues that were normal and proved to be abnormal by histologic analysis (and imaging) were not as dissimilar as might have been anticipated (nor normal and high-grade samples were the last to split into separate branches in four of the six patients). The mutation profiles for patients 1 and 3 branch early in the tree and are more distinct than that of the other patients.

We subsequently identified 77 unique mutations mapping to 29 genes across 16 chromosomes in the 23 tissue samples from the total cohort (Table E4 [online]). The majority of these mutations (53 of 77) were nonsynonymous coding mutations across 26 different genes. Additionally, a frameshift mutation in BCR was detected in samples from patients 2, 3, and 6. A frameshift mutation in TP53 was detected in the high-grade samples and histologically normal microenvironment samples that were also moderately suspicious for cancer from patient 1. There were three-stop gain mutations in EWSR1 (patients 1, 4, and 5). Hierarchical clustering of the mutations across all 23 samples was then performed to assess any underlying association or natural grouping of these mutations across the different tissue samples (Fig 4). Interestingly, the clustered subgroups corresponded to the individual patients. Six mutations across five genes (ALK, ERG, EML4, PLAU, and RET) were detected in all of the samples, with subgroups of mutations driving clusters specific to each individual (Fig 4; Tables E4, E5 [online]).

Imaging–confirmed intermediate suspicion, 31 mutations per sample ± 15; and high grade, 33 mutations per sample ± 18; Kruskal-Wallis test, \( P = .30 \). Interestingly, mutations in cancer-associated genes were observed in all of the regions sampled (including regions in which the histologic structure appeared normal [referred to here as histologically normal]) across all of the patients for the cumulative set of mutations and the filtered cancer gene mutations.

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To further delineate grouping of the samples on the basis of mutation characteristics, principal component analysis was performed (Fig 6). The first three principal components accounted for 65% of the variation of
the mutations across all tissue samples. Classification of the samples by patient and Gleason score by using the first three principal components resulted in simultaneous classification of individual patients and Gleason scores of 3+4 versus 4+5 lesions for 22 of the 23 samples. The second and third principal components only accounted for 24.9% of the mutational variance across all samples; however, by using only these two components, 20 of 23 of the individual patient samples with Gleason scores of 3+4 versus 4+5 were successfully classified (Fig E5 [online]). This separation is driven primarily by the third principal component; the genes with the largest weightings included KLK2 (rs198977), KRAS (rs712), SPINK1 (rs11319), BRCA1 (rs1799949), and BCR.

No statistically significant linear correlation was identified between individual mutations and functional MR imaging parameters, although it was noted that before multiple hypotheses corrections there was a significant correlation between two genes with contrast-agent enhancement (BRCA2, rs144848, P = .045; and ALK, rs1881421, P = .046). No significant correlations were identified between the composite PI-RADS scores and individual mutations.

Discussion

This study provides a detailed spatial radiogenomic characterization of prostate glands in six patients with confirmed high-grade tumors through evaluation of the histologically normal and abnormal microenvironments of the glands by using multiparametric MR imaging, histopathologic assessment, and whole-exome sequencing. The investigation was enabled by the technologic advancements that have been made in multiple fields, including whole-mount tissue preservation, multiparametric MR imaging capabilities, and next-generation sequencing technologies. This integrative study revealed the presence of cancer-related mutations within multiple microenvironments of prostate glands, including histologically normal and abnormal (ie, high-grade) regions; thus, exome sequencing can potentially provide a complementary role to multiparametric MR imaging and histopathology in characterizing prostate adenocarcinoma.

We observed that the background mutational spectrum in the prostate gland is greater than expected and, not surprisingly, that abnormalities found via histopathologic analysis may not be the earliest harbingers of disease. This observation supports a hypothesis that tumorigenesis in the prostate is not a discrete event, but rather part of a continuum in which unifocal versus multifocal disease are potentially within the same spectrum, in which the latter declares itself because of either diagnosis at a later stage of the disease process or a propensity for forming multiple nodules earlier because of underlying biologic variation. In light of the evidence of so-called field effects in the prostate gland and further evidence that revealed shared mutational variation in histologically normal and abnormal regions of prostates with foci of adenocarcinoma (28,29), multifocal disease may be seen as more of a distal part of the spectrum of prostate cancer from the perspective of mutational burden (20).

There is literature (30–34) that supports evidence of abnormal mutations in normal-appearing tissues in other organs including skin, lung, and ovaries. Regions that harbor abnormal mutations may presage organizational changes on the molecular level that are not yet discernable via conventional light microscopy. Changes in multiparametric MR imaging characteristics (ie, diffusion, T2-weighted signal, and tissue perfusion) reflect alterations in tissue organization, structure, and

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**Figure 5**: Phylogenetic tree generated from the mutations across all patient samples show branching and grouping according to cancer-related mutations shared within patients (vs mutations in tissue samples shown via histologic analysis to be similar). The root of the phylogenetic tree (most commonly shared mutations appears) are on the far left and the leaves of the tree (most distinct or unique mutations) are on the far right. Reading the plot from left to right, the most similar tissue samples on the basis of mutations in genes related to cancer will branch later (more toward the right). There is a branching structure of the mutations, grouping according to the individual prostate glands (including normal and histologically abnormal tissue), and separation between histologically normal versus high-grade specimens in the most terminal branches, which reflects greater similarity within each gland rather than between different glands, regardless of histologic analysis (ie, tissue samples that are histologically normal as well as abnormal from the same patient have more in common than Gleason 3+4 and 4+5 nodules from different patients). Relatively early branching of patients 1 and 3 suggest that their mutation profiles are more distinct than that of the other patients.
vascularity; however, identification of associated inconsequential genomic variation versus malignancy-associated mutations remains a challenge. For the prostate, this raises important questions to be addressed in future work, including the significance of such mutations and determining the likelihood of whether such mutations will lead to a malignant nodule, and if so, which other associated factors could drive (or avoid) such transformations.

This study had limitations. First, the sample size of the study was limited; this was primarily the result of limited availability of high-quality tissue in addition to cost considerations. Whereas the principle observation suggestive of field effect for prostate cancer is compelling and supported by other observations in the literature (28,29), identification of diagnostic mutation profiles needs to be performed with larger sample cohorts in studies designed for that purpose. Second, this was a retrospective study; this was a requisite by design and the underlying evaluated hypothesis, which required confirmed diagnoses of prostate adenocarcinoma. Although Gleason score could be assessed for the different samples, clinical outcomes or other phenotypic measures could not be performed; this is because of the indolent nature of the disease. Additional measurements would have been beneficial for confirming the significance of the measured mutations; however, limitations in tissue availability and cost curbed the ability to perform additional measurements. This unfortunately also limited the ability to assess the degree of causality of the mutations. Similarly, for normalization it would have been preferable to have obtained whole blood at the time of surgery (also a limitation of the retrospective nature of the study). However, by focusing on genes with associated cancer mutations (Catalogue of Somatic Mutations in Cancer database) and further focusing only on mutations that were identified in at least two or more different tissue samples, we were able to significantly increase the specificity of the targeted mutations.

In spite of these limitations, the assessment of the imaging, histopathologic analysis, and exomes across multiple microenvironments of the prostate in patients with prostate adenocarcinoma enabled characterization of the mutational background of the gland. Although we did not identify any significant correlations between the mutation profiles and multiparametric MR imaging measures, this may have been a limitation of the sample size because recent studies (35,36) investigating imaging correlates of multiparametric MR imaging and gene expression in prostate cancer show some promise for possible association maps to be constructed. Additionally, although detailed review by a genitourinary pathologist and genitourinary radiologist to

Figure 6: Principal component analysis of mutations across all 23 prostate tissue samples enabled stratification according to Gleason score and individual patient identification. The first three principal components of the individual genomic sampled regions clearly group by individual patients and are also able to classify different histopathologic grades of tumors according to the mutations. The third principle component separates the patients into higher (Gleason 4+5) and lower grade (Gleason 3+4) groups (demarcated by the intersecting plane composed of the first and second principal component axes in red), which shows histopathologic and mutation profile coherence.
ensure concordance of MR imaging and histologic specimens supported spatial concordance of the multiparametric MR imaging description and sequenced tissue, the development of objective, electronic registration of whole mounts with MR imaging (37–39) could assist in colocalizing lesions with greater accuracy, accounting for differences in nonconcordance of the axial axis in the MR versus the whole-mount sections (ie, obliquities between multiparametric MR imaging axial and whole-mount specimen slices).

Rad-path correlation is established with PI-RADS; however, with additional dimensions of measurement, including next-generation sequencing, it may be possible to move beyond current classification, toward molecular-Rad-Path correlations and so-called Rad-Path 2.0 (40). These results suggest that tumorigenesis in the prostate is part of a continuum in which abnormal mutations and imaging features that are moderately suspicious for cancer may predate histopathologic analysis–proven abnormalities that become apparent at a later time. Although there is evidence to support the use of targeted tissue biopsies when relying on histologic analysis alone (41), findings in this study and the mutational landscape, with a background of abnormal mutations in normal and abnormal (by histopathologic analysis) regions of the prostate, suggest that random, nontargeted biopsies may be useful in genomic-driven treatment decision making, although larger cohort studies are required to address this more definitively.

This work takes a further step in assessing the imaging and genomic features in prostate glands in adenocarcinoma, with the purpose of elucidating both the so-called natural course of the disease and the ways in which diagnostic tools can contribute to the detection of this disease. Multiparametric MR imaging and DNA mutation measurements are found to be complementary, in which the multiparametric MR imaging correlates strongly with histopathologic analysis, and DNA mutations from microenvironments proven via histologic analysis to be normal may intimate high–Gleason score nodules elsewhere in the gland, potentially reflecting latent, smoldering disease. Although these results are encouraging, additional larger studies are needed to further evaluate the potential significance of these observations and to further evaluate correlates between sequence variation and imaging characteristics.

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