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Prostate cancer histopathology using label-free multispectral deep-UV microscopy quantifies phenotypes of tumor aggressiveness and enables multiple diagnostic virtual stains

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Identifying prostate cancer patients that are harboring aggressive forms of prostate cancer remains a significant clinical challenge. Here we develop an approach based on multispectral deep-ultraviolet (UV) microscopy that provides novel quantitative insight into the aggressiveness and grade of this disease, thus providing a tool to help address this important challenge. We find that UV spectral signatures from endogenous molecules give rise to a phenotypical continuum that provides unique structural insight (i.e., molecular maps or “optical stains”) of thin tissue sections with subcellular (nanoscale) resolution. We show that this phenotypical continuum can also be applied as a surrogate biomarker of prostate cancer malignancy, where patients with the most aggressive tumors show a ubiquitous glandular phenotypical shift. In addition to providing several novel “optical stains” with contrast for disease, we also adapt a two-part Cycle-consistent Generative Adversarial Network to translate the label-free deep-UV images into virtual hematoxylin and eosin (H&E) stained images, thus providing multiple stains (including the gold-standard H&E) from the same unlabeled specimen. Agreement between the virtual H&E images and the H&E-stained tissue sections is evaluated by a panel of pathologists who find that the two modalities are in excellent agreement. This work has significant implications towards improving our ability to objectively quantify prostate cancer grade and aggressiveness, thus improving the management and clinical outcomes of prostate cancer patients. This same approach can also be applied broadly in other tumor types to achieve low-cost, stain-free, quantitative histopathological analysis.

Prostate cancer (PCa) is the most commonly diagnosed (non-cutaneous) cancer among men in the United States1. According to the National Institute of Health SEER report, over 3.25 million men in the US are currently living with this disease, and ≈1 in 6 men will be diagnosed with it over their lifetime2. Other estimates, however, suggest that the prevalence of PCa may actually be much higher. Studies using autopsy analyses indicate that over half of all men above the age of 50 harbor some form of PCa, increasing the estimate of the number of men living with this disease to over 20 million in the US3. This staggering prevalence makes it clear that, in a significant number of cases, PCa follows an indolent course; nevertheless, PCa is still the second leading cause of cancer death in men, with over 33,000 deaths in the US in 20204. Thus, while early and accurate PCA detection is critical, so too is the ability to objectively assess the tumors’ aggressiveness. Unfortunately, this remains a significant clinical challenge which has profound implications. On the one hand, there are a vast number of PCa patients...
haboring indolent tumors who are either (i) under a ‘watchful waiting’ category (i.e., wait and see if the cancer progresses) and have to live with the uncertainty of potentially having an unsampled aggressive tumor, or (ii) deemed high risk but are ultimately over-diagnosed and over-treated. On the other hand, there are many PCA patients harboring aggressive tumors who are at risk of being under-diagnosed and under-treated—for these patients an incorrect diagnosis may be fatal.

In the current standard of care, once a diagnosis of PCA has been established, one of the most important factors in assessing tumor aggressiveness is the Gleason score (this also largely dictates treatment course)\(^5^,\^7\). Here a histopathologist visually inspects hematoxylin and eosin (H&E) stained thin tissue sections and determines the two most common glandular/architectural patterns, which are assigned a grade from 3 to 5 (grades of 1 and 2 are not diagnosed on needle core biopsies, and are non-cancerous). While accepted as the gold standard, the Gleason score (the sum of the two grades) is qualitative and subject to intra- and inter-observer variability. Some studies have reported inter-observer agreement (kappa values) ranging from ~40 to 70%, with a significant portion of the discordant values dictating different treatment paths\(^8^–\^10\). H&E staining can also be highly variable and laboratory dependent, which undoubtedly contributes to observer variability, but also makes it challenging to extract quantitative parameters. Tissue fixation can also play a significant role in the variability of staining procedure\(^11^–\^13\). Thus, there is a significant need for novel technologies that can provide pathologists and clinicians with additional quantitative information regarding the aggressiveness of PCa, and thus prognosis for individual patients.

Advanced methods using genetic profiling, for example, provide a wealth of information but have shown limited success in predicting the prognosis of cancer patients\(^14^–\^20\). The poor predictive power may be attributed to the vast genetic heterogeneity of tumors, which makes it extremely difficult to identify a unique set of mutations that provide reliable prognostic information. Alternatively, recent efforts have shifted towards exploring phenotypical “common-denominators” to the countless genetic alterations\(^21^–\^27\). Phenotypical changes, including changes in metabolites, nuclear morphology, and nano-architecture, are more consistent across patients than the myriad of individual mutations and disrupted pathways underlying the disease, and can potentially better characterize tumors. This approach has shown very promising results for early cancer detection\(^21^–\^25,\^28,\^29\), and—to a more limited extent—assessing cancer aggressiveness\(^26^,\^27\).

In this work we introduce multispectral deep-UV microscopy as a novel tool for phenotyping PCA tissue sections, resulting in multiple virtual diagnostic stains and a unique quantitative biomarker that is predictive of disease aggressiveness. Multispectral deep-UV microscopy offers rich endogenous, label-free, molecular information of important tissue biomolecules with subcellular spatial resolution using a fast, low-cost imaging configuration\(^31^–\^36\). To define this novel biomarker (or phenotypical continuum), we couple deep-UV microscopy with an unsupervised analysis of the molecular signatures\(^37\). Importantly, we find that patients with the most aggressive forms of prostate cancer express a ubiquitous glandular phenotypical shift, even in glands that appear to be less aggressive. We further introduce multiple virtual “optical stains” (or “biochemical stains”) of tissue slides that highlight important components for disease diagnosis such as nuclei, cytoplasm, stroma, basal layer, nerves, and inflammation. The unique insight provided by the method is not available with current histological methods. In addition, we leverage recent advances in deep learning to translate our multi-spectral deep UV images into virtual H&E-stained images which show a high degree of correlation with the gold-standard H&E histopathological images of prostate tissue. These capabilities are important because they show that multiple types of “stains” can be rendered from a single unstained section, including novel diagnostics maps and the gold-standard H&E. This is potentially time and resource-wise efficient as it would obviate the need for chemical reagents, time consuming processes, multiple scans, or trained personnel. Plus, the unstained tissue sections remain available for any additional profiling (e.g., immunostaining, RNA labeling, etc.) or archiving without alterations. Results from this work have significant implications towards improving diagnosis and management of prostate cancer. Further, this same approach may be widely applicable to improve histopathological analysis in many other tissue types and diseases.

**Results**

**Deep-UV microscopy of prostate tissue sections.** Details of the multispectral deep-UV microscope are provided in the methods and materials section. Images were acquired from unlabeled fixed radical prostatectomy tissue samples, which were sliced (~5 µm thick) and mounted on quartz microscope slides. Images were acquired from histologically important regions containing structures with benign tissue, inflammation, stroma, high grade prostatic intraepithelial neoplasia (HGPIN), and glands with various grades of prostate cancer (Gleason grades 3, 4, and 5). Eighty-seven regions of interest were acquired from 15 patients. Each region was ~1 mm X 1.5 mm, acquired with a spatial resolution of ~300 nm. Multispectral images were taken at four key wavelengths, including 220 nm, 255 nm, 280 nm, and 300 nm (see Fig 1a). These spectral regions are chosen because the 255 nm and 280 nm bands correspond to the absorption peaks for nucleic acids and proteins, respectively, and the 220 nm band has contributions from many molecules including the two aforementioned\(^36\). The 300 nm band does not correspond to an absorption peak of any endogenous biomolecule, but is chosen as an indicator of tissue scattering which has been applied as a surrogate biomarker of tissue nano-architecture\(^36,\^38–\^39\).

To effectively represent the endogenous molecular tissue alterations that lead to cancer grade differentiation, the second processing method (Fig. 1f,g) uses an unsupervised content-preserving Deep Neural Network to show that the same multispectral data are sufficient to produce high quality virtual H&E...
stained images which is the gold-standard in pathology. The details of the Neural network are explained in the next section.

In the first method, the multispectral data were processed using a geometrical representation of principal component analysis (PCA), which is an unsupervised method. In this process, approximately 130 million spectra from select representative regions were used to calculate the principal components (PCs). Figure 1b shows the resulting orthogonal PCs. It is important to note that these vectors, while purely mathematical in nature, in fact resemble the absorption and scattering spectral behavior of biological tissues. For example, the first principal component, PC1, shows a unipolar, monotonically decreasing behavior that is consistent with the expected response of tissue scattering. PC2 and PC4 show peak responses that correspond to protein absorption, while PC3 shows an inverted peak that is in agreement with the absorption from nucleic acid. Nevertheless, projections of the spectra onto these PCs do not uniquely correspond to these molecules, and do not prominently highlight important tissue structures alone, as seen in Fig 1c.

To obtain a more natural representation of the endogenous tissue composition, we transform these data from a Cartesian coordinate system with only the first three PCs (which possess over 99% of the data variance) to spherical coordinates (see Fig 1d). (The same procedure can be applied with any combination of three PCs.) In this representation, the azimuth (θ) and elevation (φ) angles contain all the information about the shape of the spectra; in other words, these two dimensions contain nearly all the available biophysical and biochemical

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**Figure 1.** Processing of multi-spectral deep UV images using the two proposed methods. Top: geometrical representation of principal component analysis. Bottom: Unsupervised content-preserving transformation for optical microscopy. (a) A representative multi-spectral deep UV transmission data cube taken at 220, 255, 280 and 300 nm. (b) The 4 principal components resulting from 130 million spectra from representative select regions. (c) Calculated projections of the data cube on the principal components. (d) Schematic of data conversion from Cartesian to Spherical coordinates. (e) Representative 2D histogram of a data cube using azimuthal and elevation coordinates. (f) Schematic of trained Neural network. (g) Representative virtually H&E stained prostate tissue (Network output).
information. The radius, on other hand, serves as a relative measure of the concentration. Thus, images can be represented in a hue-saturation-value (HSV) color space, with the hue given by the angular coordinates (either elevation or azimuth angle, as shown in Fig. 1e), the value set by the radius, and the saturation fixed to 1.

An important feature of this geometrical PCA representation is that each point in the 2D histogram of the elevation and azimuth angles (Fig. 1e) represents a unique spectral response, and hence different molecular and/or biophysical makeup. Thus, by color-coding images based on the angular distributions, we are able to assign a unique hue to spatial regions with similar composition. The resulting “optical stains” enhance contrast among various structures in prostate tissue sections which can be leveraged, along with H&E, for diagnostic applications.

Figure 2 shows two types of “optical stains” that highlight important tissue structures. In the first (Fig. 2a), the elevation angle is used to encode hue which yields the most prominent contrast for cell nuclei, depicted in green. This is consistent with the general behavior of the PCs, as the elevation angle in this case corresponds to a ratio of the 3rd PC (which resembles the inverted absorption peak from nucleic acids) relative to both the
Prostate cancer diagnosis and grading using deep-UV microscopy. Using a 3D space defined by the first three PCs, we defined a third “optical stain” by encoding hue using the azimuth angle (Fig. 4). These maps highlight contributions from light scattering (from PC1) relative to both proteins and nucleic acid (PC2 and PC3, respectively). We note that scattering variations arise from genetic and epigenetic perturbations that result in micro and/or nano-scale alterations in intracellular milieu, such as the cytoskeleton, ribosomes, chromatins, mitochondria, and collagen fibrils that are known to be altered in field carcinogenesis. Furthermore, protein and nucleic acid alterations have also been well documented throughout the progression of prostate cancer.

The resulting image representation (i.e., optical stain) does not exhibit contrast to structures conventionally used in histopathology (e.g., nuclei, cytoplasm, stroma, etc.); instead, we find that this representation encodes for a glandular phenotype that correlated with malignancy. Figure 4 shows two examples from patients with intermediate-grade cancer. Here benign glands possess a blue hue, while glands with cancer (Gleason grades 3 or 4) exhibit a relative shift captured in green to red hues which represents an increase in nucleic acid and protein intermediate-grade cancer. Here benign glands possess a blue hue, while glands with cancer (Gleason grades 3 or 4) exhibit a relative shift captured in green to red hues which represents an increase in nucleic acid and protein content, potentially from cell overgrowth byproducts. In these maps, the glands were segmented for clarity (performed manually for simplicity here, though this process can be automated). Again, the change in color represents alterations in the scattering properties relative to protein and nucleic acid content, all of which have been implicated in early-stage alterations of cancer, as well as metastatic disease. Thus, the azimuth angle from a geometrical representation of the first three PCs effectively yields a phenotypical continuum that can be applied as a surrogate biomarker of prostate cancer malignancy.

It is worth highlighting important features in Fig. 4. Figure 4a,b show a set of pseudo-neoplastic benign glands (blue arrows) that are not well formed, meaning they express slight cytological and morphological variations such as cytoplasm clearing that classifies them as a mimicker of prostatic adenocarcinoma (typically of Gleason grade 3). However, the existence of basal cells around the glands as well as the papillary infoldings of the gland differentiates them from carcinoma. And indeed, the malignancy optical stain clearly indicates that these glands are benign and distinct from Gleason grade 3 and 4 glands (green and red arrows, respectively). Figure 4c,d show benign central zone histology glands (blue arrows) surrounded by benign glands (green arrows). Central zone histology glands are potential mimickers of HGPIN and Gleason Grade 4 cancer glands (Cribiform) and are often difficult to differentiate from cancer glands; nevertheless, the malignancy optical stain identifies these glands as benign. Further, Fig. 4d clearly shows a gradual color gradient, and hence phenotypical continuum, from left to right as the glands progress from benign to cancer. It is important to note that even though third optical stain shows a higher degree color variability/noise, the general meso-scale behavior clearly correlates with malignancy. It is clear that the information provided by this optical stain is independent and complimentary to the gold standard H&E stain.

Supplemental Fig. S2 shows additional examples from patients with aggressive disease (i.e., those containing Gleason grade 5 glands). These samples possess a unique response which is discussed in more detail below.

To investigate the properties of this phenotypical shift further, we analyze the cumulative behavior of benign glands and cancerous glands with the same Gleason grade for each patient. In this process, cumulative 2D histograms were generated for each type of gland (benign and Gleason grades 3-5) for each patient, then data were integrated across elevation angle, and finally the center of mass (CoM) of the resulting azimuth angle distributions were computed. This value effectively quantifies the hues in the malignancy optical stain shown in Fig. 4. Figure 5 shows the results, with Fig. 5a showing the absolute azimuthal CoM for all the benign glands for each patient. This value is then taken as a basis for all other (cancerous) gland types for each patient, thus providing a personalized reference point for a malignancy biomarker.

Figure 5b,c show the relative shifts in the CoM of cancerous glands relative to the benign glands of each patient (absolute shifts are shown in Fig. S3). Self-calibration with respect to benign gland of each patient is necessary to reduce significant inter-patient variability which can be very large and, if unaccounted, can potentially obscure signals of interest. A remarkable result of this personalized biomarker is that patients with the most aggressive form of prostate cancer (i.e., those containing Gleason grade 5 glands) exhibit a ubiquitous glandular phenotypical shift in the opposite direction as patients with less aggressive forms of prostate cancer. That is, for...
Figure 3. Comparison of the two “optical stains” with corresponding H&E-stained tissue scans from various prostate tissue structures. (a) Benign gland. (b) High grade Prostatic intraepithelial neoplasia (PIN) region. (c) Cancer region with Gleason Grade 3 glands. (d) Cancer region with Cribriform Gleason Grade 4 region. (e) Region with Gleason Grade 5. (f) Region with necrosis inside a Gleason Grade 5 cancer gland. Necrosis is clearly distinguishable from the cancer cells on the left side of the image. (g) Inflammation. (h) Red blood cells.
patients with aggressive cancer, both Gleason grade 5 regions and lower Gleason grade regions (3 and 4) show a negative azimuth CoM shift, which is the opposite behavior compared to patients with less aggressive tumors. This unique and ubiquitous shift—only present in aggressive prostate cancer—may be attributed to higher contributions from scattering which is indicative of changes in tissue organization at the nanoscale level. Similar behavior has been reported in other studies of the scattering properties of cancerous tissues\cite{37,38,40,41} and is likely related to the field effect of carcinogenesis.

Figure 4. Comparison of malignancy maps from different prostate regions with corresponding H&E scans. Insets show the 2D histograms for comparison. As clearly evident from Fig. 4 the malignancy optical stain shows diagnostic capabilities complimentary to H&E, where only morphological parameters are considered. In (b and d) we have manually removed stroma and inflammation regions to aid visibility. The small-size red regions on the edges of the glands are originating from two sources: 1- ill-formed fused type Gleason Grade 4 regions have spread around benign glands and in fact it is an indicative of existence of cancer. These regions are sometimes missed by pathologists in H&E analysis 2- Existence of inflammation cells that were impossible to remove with manual segmentation. These inflammation cells are limited in number and do not contain any diagnostically important information and they are mostly in stromal region. (Scale bar: 200 μm).
UTOM for label-free H&E colorization with UV microscopy. The novel optical stains presented above provide unique insight into tissue structures based on endogenous molecular composition, nanoscale structures, and PCa aggressiveness; nevertheless, H&E-contrast is imperative for PCa diagnosis and grading. While additional tissue can be stained with H&E, it is also possible to translate the UV images into virtual H&E images to enable visualization of the same exact specimens, down to the subcellular level, in different diagnostic formats/stains. This label-free pipeline also avoids cumbersome, time-consuming, and complex procedures, avoids stain artifacts and variations that are common with H&E, and finally, the original unstained tissue can be preserved for further processing or archiving.

To translate the label-free UV images into virtual H&E images, we apply a recently developed unsupervised content-preserving transformation for optical microscopy (UTOM) deep neural network. UTOM adapts the general framework of cycle-consistent generative adversarial networks (Cycle-GAN) which can transform images from one domain into another without requiring pixel-level paired data. In UTOM, a forward and backward GAN are trained simultaneously to learn a pair of opposite mappings between the UV and H&E image domains, as shown in Fig 6a. In this process, a cycle-consistency loss constraint, and a pair of saliency constraints are imposed to correct for mapping direction, which avoids distortions (Fig. 6a). In the training process, the overall
network converges when the discriminators cannot differentiate between images produced by their generators (i.e., when the two GANs reach equilibrium; see Fig. 6b). Once trained, new images can be fed into the network and transformed into the desired domain (Fig. 6c). This approach has been used for image restoration (e.g., resolution enhancement, removing distortions), for virtual fluorescence labeling of label-free phase images, and H&E virtual staining of autofluorescence images. The training set for this work comprised of 54 regions from 10 patients, while the test set (transformation group) contained 21 regions from the remaining 5 distinct patients. More details on the training process and final image translation are provided in methods and materials section.

Figure 7a–d show two representative examples of UV translated (virtual H&E) prostate tissue images from the test set, along with their corresponding adjacent H&E-stained sections. The figures clearly show that the UV translated images are nearly identical to the H&E-stained tissues sections, with the most marked differences arising from the fact that the images are from adjacent sections. Specifically, the virtual H&E images preserve or improve several important features that play an important role in PCa diagnosis: First, as shown in Figs. 7b, f, j and n the UV translated images successfully recapitulate the appearance of basal cells and basal cell lamina around benign glands which are of utmost importance for PCa diagnosis (corresponding glands on H&E image are shown in Figs. 7e, i and m for comparison). This feature is also observed in Fig. 7l where an entrapped benign gland is clearly differentiated from surrounding cancer regions (the adjacent H&E stained image is shown in Fig. 7k). Second, PCa regions shown in Figs. 7g, h, o and p depict luminal epithelial nuclei with more consistent (and arguably improved) contrast in the UV translated images compared to their corresponding H&E-stained sections. These types of structures are especially important in differentiating cancer glands from other mimickers of cancer where the structure of the gland is slightly disrupted. Finally, the appearance of the clear or pale eosinophilic cytoplasm as well as hyperchromatic nuclei are well preserved, which in some cases can be indicative of PCa Gleason Grade 4.

Virtual H&E evaluation. To assess the quality of the UV translated, virtual H&E images compared to the gold standard H&E-stained images, we conducted a panel study with 4 board-certified/board-eligible histopathologists. Here the pathologists evaluated a total of 42 large area images (~1 mm × 1.5 mm), half of the images (21) were images of H&E-stained tissue sections and the other half (21) were virtual H&E from the same
regions (adjacent slices), all from the UTOM test set. Two pathologists (group 1) were assigned a set of 21 images comprising a mixture of virtual and stained H&E images, and the other two pathologists (group 2) were assigned the complimentary set, meaning images from the same regions but switching virtual H&E images with the images of stained H&E sections, and vice versa (no pathologist viewed the same region in both the virtual H&E and H&E-stained formats). While reviewing the images, pathologists were asked a series of questions regarding the quality of the images, with numerical scores ranging from 1 (poor) to 3 (excellent). They were also asked to provide a Gleason score for each region.

Results of the panel study are summarized in Table 1. Data show that the UV translated virtual H&E images and the H&E-stained tissue section images have very similar quality as assessed by the pathologist panel. With the exception of the nucleolus quality, which was evaluated slightly lower in the virtual H&E format, all other structures were assessed to have the same quality between the two modalities, with no statistically significant differences. The gland quality, which is of particular importance for PCa diagnosis, was deemed nearly identical between the two methods, as was the cytoplasm quality. Most importantly, the pathologists’ diagnostic confidence was very similar for both methods (and not statistically different). We attribute the small difference in nucleus quality to the presence of lipid-laden macrophages (Xanthoma), mesonephric remnants, and hyperchromatic nuclei in a few regions with inflammation, which have a slightly disrupted visible quality in the translated images. However, (1) these are not diagnostically meaningful (which is likely why the diagnostic confidence remained the same between the two groups, even though the nucleus quality was slightly lower in the UV translated images), and (2) the nucleus quality can be improved with additional training.

Figure 7. Comparison of translated virtual H&E images and corresponding H&E stained scans. Examples of (b)–(d) Two predicted output virtual images (a) and (c) along with their reference H&E images. (e)–(p) show three selected zoomed regions for each area. These regions have been selected to compare features on both H&E and translated virtual H&E images.
chondria, and collagen fibrils\textsuperscript{37,38,40–43}. Along with protein and nucleic acid alterations\textsuperscript{44–49,51,53,55,56}, changes in these and/or nano-scale alterations in the intracellular milieu, such as the cytoskeleton, ribosomes, chromatin, mitochondrial scattering signatures in conjunction with proteins and nucleic acids. These maps are likely indicative of micro-structural features of the tissue that can be observed with the gold standard H&E stains, completely new structures are observed when incorporating the unique UV spectral signature combined with an unsupervised spectral analysis to transform the multi-spectral data cubes into phenotypical maps or “optical stains” with subcellular spatial resolution. The spectral analysis suggests that the main contributing factors to these maps arise from scattering which serves as an indicator of tissue nano-architecture, and from proteins and nucleic acids. However, we do not rule out contributions from other molecules\textsuperscript{64–68}. Maps derived primarily from spectral signatures that correlate with proteins and nucleic acids provide high contrast among various critical tissue components, including nuclei, cytoplasm, basal layer, stroma, and glandular tissue, which can enhance our ability to recognize anomalies in prostate tissues.

In this study, we have introduced multi-spectral deep UV microscopy as a novel, fast and reliable method to capture quantitative molecular and nano-scale information from unlabeled prostate tissue sections. We have utilized the unique UV spectral signature combined with an unsupervised spectral analysis to transform the multi-spectral deep UV microscopy data into high quality virtual H&E images, with a diagnostic accuracy statistically equivalent to the gold standard. We also calculate the inter-group concordance for grade group decisions using the H&E and virtual H&E images. The results (Table 1) show inter-observer variability at similar levels to what has been reported in previous studies\textsuperscript{64–68}. Importantly, however, the concordance in Gleason grade decisions is very similar between the UV translated virtual H&E images and the H&E-stained tissue section images within each group. These results strongly suggest that the format of the images (virtual H&E and H&E stained) did not play a role in the concordance levels. It is also worth noting that the agreement between the two most senior, board-certified pathologists was very high—they agreed in 17 out 21 regions even though they were viewing each region in different formats (one was in group 1 and the other in group 2). The 4 regions of discordance were between boarder line Gleason grades 3 and 4.

Finally, we calculate the accuracy of the Gleason scores provided by the pathologists for both the H&E and virtual H&E images. For this task, we first select a “ground truth” given by the decision of one of the two senior board-certified pathologists (whichever one of the two whose decision was based on the stained H&E section for a particular region was selected). Results show that the accuracy of the Gleason grades using H&E and virtual H&E images are similar (i.e., not statistically significant), with 72.5% accuracy for H&E and a slightly higher 77.45% accuracy for virtual H&E (p-value = 0.24). Alternatively, only using the regions where both senior, board-certified pathologists agree (17 out 21) and using their assessment as “ground truth”, we find an accuracy for Gleason grade of 73.6% for H&E and 81.6% for the UV translated, virtual H&E images (again, differences not statistically significant, p-value=0.42). This is a more robust “ground truth” but omits any data lacking concordance from the two senior pathologists. Nevertheless, regardless of the selected “ground truth”, the grading accuracy values show that the proposed UTOM method is capable of transforming the multispectral deep UV data into high quality virtual H&E images, with a diagnostic accuracy statistically equivalent to the gold standard.

**Discussion**

In this study, we have introduced multi-spectral deep UV microscopy as a novel, fast and reliable method to capture quantitative molecular and nano-scale information from unlabeled prostate tissue sections. We have utilized the unique UV spectral signature combined with an unsupervised spectral analysis to transform the multi-spectral data cubes into phenotypical maps or “optical stains” with subcellular spatial resolution. The spectral analysis suggests that the main contributing factors to these maps arise from scattering which serves as an indicator of tissue nano-architecture, and from proteins and nucleic acids. However, we do not rule out contributions from other molecules\textsuperscript{64–68}. Maps derived primarily from spectral signatures that correlate with proteins and nucleic acids provide high contrast among various critical tissue components, including nuclei, cytoplasm, basal layer, stroma, and glandular tissue, which can enhance our ability to recognize anomalies in prostate tissues.

While the “optical stains” derived from proteins and nucleic acids correlate well with the overall structures observed with the gold standard H&E stains, completely new structures are observed when incorporating the scattering signatures in conjunction with proteins and nucleic acids. These maps are likely indicative of micro- and/or nano-scale alterations in the intracellular milieu, such as the cytoskeleton, ribosomes, chromatin, mitochondria, and collagen fibrils\textsuperscript{47,48,49,50,51,52,53,54,55,56}. Along with protein and nucleic acid alterations\textsuperscript{44–49,51,53,55,56}, changes in these structures have been implicated in the field effect of carcinogenesis. Indeed, here we observe that these structures map benign glands to different hues compared to cancerous glands, effectively yielding a “malignancy map.” By quantifying these relative phenotypical shifts, we also find that cancer patients with the most aggressive forms of prostate cancer (those with Gleason grade 5 glands) possess a ubiquitous and unique phenotypical shift compared to patients with less aggressive cancers.

These results have significant implications. Because less aggressive cancer glands (e.g., Gleason grade 3) possess a different phenotypical shift in patient harboring an aggressive cancer (those with Gleason grade 5 glands), this phenotype or biomarker may help identify patients with aggressive forms of prostate cancer even if initial biopsies miss the more aggressive regions. These results could have profound implications for the analysis of random prostate tissue biopsies which cannot cover the entire organ and are hence susceptible to missing cancer regions. It is worth emphasizing that this is achieved by defining a continuous quantitative marker, that evaluates the malignancy level of each gland with respect to benign glands of the same patient, and avoids the use of Gleason grade (though we use Gleason grade to establish a correlation to this accepted standard).

While this continuous biomarker does not show appreciable differences between Gleason grade glands 3 and 4, incorporating morphological features along with this biomarker may potentially improve our ability in identifying/grading prostate cancer. For instance, quantitative information from the UV spectra and derived optical maps can help differentiate anomalous benign glands that mimic cancer and can be difficult to detect. Furthermore, all the images, supported by their histograms of the molecular signatures, show that healthy tissue,

| Parameter                              | H&E   | Virtual H&E | Statistical significance |
|----------------------------------------|-------|-------------|--------------------------|
| Nucleus quality (scale of 3)           | 2.57  | 2.32        | *                        |
| Cytoplasm quality (scale of 3)        | 2.45  | 2.42        | N.S                      |
| Gland quality (scale of 3)            | 2.60  | 2.57        | N.S                      |
| Diagnosis confidence (scale of 3)     | 2.26  | 2.16        | N.S                      |
| Inter-group concordance for group 1   | 81.82%| 85%         | -                        |
| Inter-group concordance for group 2   | 60%   | 50%         | -                        |

Table 1. Comparison of translated UV images with H&E. N.S.: Not statistically significant. *p < 0.05.
disease regions and their underlying composition span a continuum rather than a discrete distribution. This is in line with our understanding of disease progression and may help better characterize prostate cancer compared to discrete labels (as with Gleason grades). This new information may also help assess a more ideal personalized treatment course for patients.

These results, while from a small sample size, lead to more fundamental questions: Do patients need to have this unique malignant phenotype to develop the aggressive form of PCA? If so, can it be detected even before aggressive cancer develops? How early? Or is there a ubiquitous switch across the gland that occurs once the disease progresses to this more aggressive form? The answer to these questions requires further understanding of this malignant phenotype and will guide our future work as a larger sample size is analyzed.

Finally, using a state-of-the-art deep learning algorithm, UTOM, we showed that the UV images can be readily translated into virtual H&E images that accurately mimic the structures and colors present in the gold standard bright-field microscopy images of H&E-stained prostate tissue sections. This process is advantageous from a histopathology viewpoint because multiple diagnostic (virtual) stains can be produces from the same exact regions. The process also avoids the need for laborious, time-consuming, and costly chemical staining procedures, avoids staining viability, and preserved tissue for other uses. A panel of board-certified/board-eligible pathologists assessed the quality and diagnostic potential of the UV translated images to be equivalent to the gold-standard H&E-stained tissue section images.

Furthermore, other optical technologies have shown very promising results to help improve histopathology, using methods such as UV excited florescence/auto-florescence, infrared, and Raman Scattering. However, there are important limitations associated with each approach. For instance, infrared imaging technologies provide rich molecular information, but have complex and expensive equipment, are relatively slow, and lack critical subcellular and cellular level resolution. UV excited florescence methods have demonstrated rapid visualization of subcellular H&E level histology in thin or thick tissues and have garnered a lot of attention. However, these methods are not quantitative, often require exogenous agents, and to our knowledge have not been shown to provide novel diagnostic information. Similarly, auto-florescence based methods have been encouraged to generate label-free H&E-like images but auto-florescence intensity differs from patient to patient, the signal to noise ratio is low, and level of endogenous molecular contrast is limited, all of which increases uncertainty and hence the number of misdiagnosed disease cases. Raman microscopy/spectroscopy methods provide rich molecular content and allow differentiation of malignant tissue. However, Raman scattering is a weak process that requires long acquisition times and signal can easily be obscured by fluorescence. Nonlinear coherent Raman imaging is much faster but systems are complex and expensive. Finally, most of these optical technologies suffer from complexities in how to integrate into current pathology practice workflows.

The label-free multi-spectral deep UV microscopy approach proposed here shows unique capabilities which overcomes many of the limitations of other methods described above or which can be complimentary to help improve diagnosis and grading of prostate cancer. This approach is high-resolution (~300 nm) (which provides approximately two times or more better resolution than standard H&E practice and commercial digital scanners), provides rich molecular and nanoscale quantitative information, and is simple and low-cost (~$20k; and the UV transparent quartz slides used here could also be replaced by cheap UV transparent polymers). The approach is also widefield with exposures of ~100ms per field of view (~170 μm x 230 μm) making it relatively fast. The estimated total acquisition time for all 4 wavelengths with our current system for a 10 mm x 10 mm slide is ~240 seconds which yields a throughput of 15 slides/hr (for reference, commercial digital pathology scanners have a throughput of 20–80 slides/hr). With further improvements in the setup such as automated scanning algorithm and auto-focusing, and/or using lower resolution/magnification to match standard practice (which would yield much larger fields of view per acquisition), much higher scanning rates are achievable. Such fast scanning rates suggest that our proposed deep UV microscopy method is quite feasible for routine clinical practice and potentially surgical pathology. This approach could also be combined with other state-of-the-art structure-based neural networks recently introduced to help automate diagnosis.

In conclusion, we have introduced label-free multispectral deep UV microscopy to help analyze prostate cancer histopathology. We have demonstrated the unique capabilities of this method, which can help improve diagnosis and management of prostate cancer. Finally, this same quantitative approach can be applied broadly across histopathological analysis of many tissue types and diseases. To our knowledge, this is the first demonstration of the utility of transmission-based deep UV microscopy for the analysis of tissue sections and histopathology. Future work will focus on imaging and analyzing a larger sample size (prostate and other tissue types) to further show the robustness of this method.

Materials and methods

Deep-UV multispectral microscopy set up. The deep UV transmission images were obtained using a microscope system that consists of a plasma-driven broadband light source (Energetiq, EQ-99X) that provides a continuous spectrum from 200 nm to 2 μm. The output light from the source is focused on the sample using an off-axis parabolic mirror (Newport). A long-pass dichroic mirror is used to filter out the wavelengths of light above ~550 nm. For each region of interest, a multispectral data cube is captured using bandpass filters (bandwidth = 10 nm) centered at 220, 255, 280 and 300 nm. The filters are placed on a filter wheel to change the imaging wavelength of the system. A 0.5 N.A. UV objective (Thorlabs LMU-40X-UVB) is used to collect the transmitted light and a biconvex (f = 150 mm) lens is used to relay light onto a UV camera (PCO. Ultraviolet). A schematic of the setup is shown in Fig. S1. For each acquisition, the camera integration time was ~100 ms. Each captured region of interest represents a field of view of about ~170 μm x 230 μm. The resolution of our system is ~300 nm. In this work, we studied regions that were comprised of 64 tiles in the form of an 8 by 8 mosaic.
image. To enable reliable stitching, each tile has ~15% overlap with its neighbors. The final resulting region is approximately ~1 mm × 1.5 mm.

**Sample collection and preparation.** Paraffin-embedded formalin-fixed blocks from radical prostatectomy specimens were obtained from 15 prostate cancer patients. All the patients had not received any neoadjuvant therapy prior to radical prostatectomy. The Gleason scores (Grade groups) and tumor stages were assigned by Urologic Pathologists in all cases. Next thick slices (~5 microns thick) of the tissue blocks were mounted on quartz slides and were deparaffinized by incubating the slides in Xylene bath for 5 minutes. The samples were then placed in 95% Ethanol for 3 minutes to remove Xylene and washed with dionized water. One section was used for UV imaging and a second section was stained with H&E and imaged with a bright field microscope.

All tissues are de-identified from archived tissue block for Emory University Hospital (n = 10) or a commercial vendor (Biomax) (n = 5). The Institutional Review Board of Georgia Institute of Technology reviewed and approved all protocols (H16343 protocol). Informed consent was obtained from all patinets and/or their legal guardian(s). All methods were carried out in accordance with relevant guidelines and regulations.

**Data processing.** To study the molecular content of the imaged tissue slides, different wavelengths in each captured multispectral data cube were registered in MATLAB (Mathworks) Environment. Next, in order to have a single wide-field UV image we used an image stitching code (MIST)82, developed by National Institute of Standards to stitch the 64 tiles captured separately.

To calculate the principal components (PCs) of the multispectral prostate tissue images, we selected 90 regions that yielded approximately ~130 million spectra which represented all biologically important structures in prostate tissue. Next, we performed PCA in MATLAB to calculate the 4 principal components of the selected regions.

To generate color-coded images, we calculated the projections of the multispectral UV data on PC 1, 2, 3, and 4, respectively. Next, we converted the resulting projection vectors (Proj 1, Proj 2, Proj 3) and (Proj 2, Proj 3, Proj 4) from Cartesian coordinates to Spherical coordinates (Azimuth (θ), Elevation (ϕ), Radius (R)), where Proj 1 represents the projection of UV data on PC1. Finally, to get the geometrical representation of the PCA, we calculated a two-dimensional histogram of the azimuth (θ) and elevation (ϕ) angles for each case. Lastly, colorized the images using a Hue-Saturation-Value (HSV) color space, where the hues are assigned based on either azimuth or elevation angle, the value is set by the radius and the saturation is set to 1.

**Calculation of the azimuthal shifts.** To calculate the azimuthal shifts that are correlated with prostate cancer grades, first we annotated all the corresponding H&E images with appropriate Gleason grades. The annotations were reviewed and approved by a board-certified Urologic pathologist. Next, for each patient, the multispectral UV data were manually segmented according to the approved H&E annotations to extract all the pixel spectra that have the same Gleason grade. Once all the grade specified spectra were collected, we calculated cumulative 2D histograms using Principal components 1, 2, and 3 for each Gleason grade category as described in the data processing section. Finally, we integrated each 2D histogram in elevation direction to generate the azimuth dependent graph of molecular content, and recorded the Azimuth coordinates center of mass. We repeated this procedure for all the captured regions from all the patients.

**Virtual H&E Colorization using UV microscopy images.** To perform machine learning process, we used the label-free UV images of the unstained tissue sections from 15 patients from all 4 wavelengths (220, 255, 280 and 300 nm). For each captured region the corresponding H&E-stained image from adjacent slice were used as a reference. All the UV and H&E images were scaled to the same pixel size (90 nm). Next, we used 54 regions from 10 patients that contained representative biologically structures in prostate tissue, as the training data-set for our model (~13.5 billion spectra). The remaining regions (21) from the other 5 patients were used as the testing data set to evaluate the color transformation model. The important point about the testing data set is that the regions come from completely independent patients and no regions from testing patients are used in the training process. In the training dataset, the 4-channel UV data and H&E images (RGB channels) were randomly cropped into 512 × 512 patches. The total numbers of UV and H&E patches are 64,336 and 81,667, respectively (Fig. 6a). During the test phase, the UV images were first partitioned into small patches with 25% overlaps. After a model was trained, patches from the previously unseen 5 patients were then fed into the model to generate the corresponding H&E patches. To finally form a large area virtual H&E image (each ~1 mm × 1.5 mm), we cut out the boundaries (half of the overlap) of the generated patches and stitched the remaining parts together one by one.

**Virtual H&E Color normalization.** To remove undesirable color variations of the H&E-stained histological images, which result from differences in staining protocols, slide scanners and other factors, we adopted the structure-preserving color normalization (SPCN) method proposed by Vahadane et al.89. For a given image, we first estimated its stain density maps and color appearances via sparse non-negative matrix factorization. Then, we combined the stain density maps with a stain color basis of an arbitrary target H&E image so as to change only the color appearances while preserving the structure of the source image.

**UTOM method.** To produce virtual H&E colorized images, a forward GAN and a backward GAN are trained simultaneously to learn a pair of opposite mappings between two image domains. Along with the cycle-consistency loss, a saliency constraint is imposed to correct the mapping direction and avoid distortions of the image content. For each domain, a discriminator is trained to judge whether an image is generated by the genera-
tor or from the target domain (Fig. 6b). When the loss converges, the two GANs reach their equilibriums, which means that the discriminators cannot distinguish images produced by their generators from the target images. An image could be mapped back to itself through the sequential processing of the two generators, and more importantly for biomedical images, the saliency map keeps high fidelity after each transformation (Fig. 6a). The well-trained generator G of the forward GAN is used for transformation task from UV images to H&E images (Fig. 6c).

The architectures of the generator and the discriminator are visualized in Fig. 6b. The first three layers of the generator are downsampling layers implemented by strided convolution to extract low-level abstract representations. Nine stacked residual blocks are followed to extract high-level features. The number of residual blocks reflects the model capacity. More residual blocks are recommended for more complex tasks. It is important to note that to reduce the training cost and obtain a comparably good performance, we used the Residual block generator instead of U-Net. The residual block design can also alleviate the problem of vanishing/exploding gradients when deeper networks are adopted, and converge much faster than standard solvers. The last three upsampling layers are also implemented by strided convolution. They are used to integrate extracted features and rescale the image to its original size. The discriminator is a relatively shallow CNN. Each layer downsamples the feature maps but doubles the channel number. The last convolution layer generates a single-channel feature map and classification is performed on each element of this feature map (PatchGAN classifier). The final true or false label is generated by averaging individual labels of all elements. Each convolution layer in both the generator and the discriminator contains a nonlinear activation unit. Whether to use the sigmoid function or rectified linear unit (ReLU) is marked with corresponding arrows in Fig. 6b.

The Adam optimizer was used to optimize network parameters. The initial learning rate is 0.0002, which decays linearly every 50 iterations with a rate of 0.99. The batch size was set to 1 and the images were flipped randomly for data augmentation. We trained the network for about 5 epochs, with about 80000 iterations in each epoch. On a single NVIDIA GEFORCE RTX 2080 Ti GPU (11GB memory), the whole training process took approximately 48h. After training, UTTOM took 21ms to generate a $512 \times 512$ H&E patch and cost 3s to produce a whole-slide HE image.

We used a PC system with an Ubuntu 16.04 LTS operating system and a CPU Intel(R) Xeon(R) CPU E5-2683 processing unit. Also a PyTorch 1.6 was used as the Deep Learning Framework and Python 3.7 for image processing.

**Virtual H&E evaluation methodology.** We prepared a web-based survey including 21 unidentified, mixed H&E and virtual H&E regions (group 1, 10 H&E and 11 virtual H&E and group 2, 11 H&E and 10 virtual H&E images of the same regions) and asked 2 board-certified and 2 board-eligible pathologists to submit their evaluations of the quality of parameters such as nucleus, cytoplasm and gland quality. Further, we asked them to submit a Gleason Score for each region to compare the accuracy of diagnosis for both H&E and virtual H&E images. Each question was based on the scale of 1 to 3 (1 for poor, 2 for moderate and 3 for very good quality). The responses were downloaded and used for statistical analysis. This clinical panel review protocol (no. H19389) was Institutional Review Board-exempt.

**Data availability**

Additional Colorized Images along with the corresponding H&E references are available at: https://zenodo.org/record/5140334.

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Author contributions
S.S. contributed to the experimental design, performed experiments and data acquisition, performed data processing and computational data analysis, histopathological analysis and evaluation and tissue annotation, wrote the manuscript and prepared the figures. A.O contributed to the experimental design and data acquisition. H.Q. developed the machine learning model and performed deep learning image translations. N.K. contributed in data acquisition. X.L. contributed to machine learning modeling. A.O.O. was the histopathology supervisor, approved prostate tissue annotations and contributed to translated UV image evaluation. Q.D. supervised the machine learning processes. F.E.R. was project supervisor (Principal investigator), designed experiments, wrote the manuscript, and supervised all steps. All the authors contributed to final approval of the manuscript. F.E.R. and S.S. performed manuscript submission and review process.

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Competing interests
The authors declare no competing interests.

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