Co-Registration of ex vivo Surgical Histopathology and in vivo T2 weighted MRI of the Prostate via multi-scale spectral embedding representation

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Multi-modal image co-registration via optimizing mutual information (MI) is based on the assumption that intensity distributions of multi-modal images follow a consistent relationship. However, images with a substantial difference in appearance violate this assumption, thus MI directly based on image intensity alone may be inadequate to drive similarity based co-registration. To address this issue, we introduce a novel approach for multi-modal co-registration called Multi-scale Spectral Embedding Registration (MSERg). MSERg involves the construction of multi-scale spectral embedding (SE) representations from multimodal images via texture feature extraction, scale selection, independent component analysis (ICA) and SE to create orthogonal representations that decrease the dissimilarity between the fixed and moving images to facilitate better co-registration. To validate the MSERg method, we aligned 45 pairs of in vivo prostate MRI and corresponding ex vivo histopathology images. The dataset was split into a learning set and a testing set. In the learning set, length scales of $5 \times 5$, $7 \times 7$ and $17 \times 17$ were selected. In the independent testing set, we compared MSERg with intensity-based registration, multi-attribute combined mutual information (MACMI) registration and scale-invariant feature transform (SIFT) flow registration. Our results suggest that multi-scale SE representations generated by MSERg are found to be more appropriate for radiology-pathology co-registration.

In spite of different radiographic imaging modalities (e.g. MRI, Ultrasound) being available for prostate cancer diagnosis, a definitive ascertainment of disease extent is only possible by histopathologic examination on surgically excised specimens. Diagnosis and delineation of prostate cancer regions based on routine MRI may suffer from inter-rater differences. Figure 1 illustrates the challenges of relying solely on radiology readers to determine ground truth for disease extent using imaging alone.

Consequently, there has been a recent appreciation of the need to identify alternative image representations that can complement intensity information which includes image gradients, co-occurrence information, and image segmentations for the purposes of multi-modal image co-registration. Texture features, e.g. Gabor wavelets provide multi-scale, multi-oriented textured representations of the original images. The intuition behind the texture feature representation is that similar underlying structural attributes (e.g. edge patterns) can be identified in the two images to be registered. Texture features extracted at differently sized neighborhoods enable the capturing of different underlying structural cues. In addition, certain scales might result in expression of cues or features that are not expressed at other scales. Hence, it becomes critical to explicitly consider the scales at which the features are extracted and develop better representations in a scale-specific manner. The challenge though is...
Identifying which representations are most appropriate for multi-modal co-registration. Previous registration methods involving texture analysis did not explicitly interrogate the role of texture as a function of feature scale. MSERg aims to leverage the use of multiple feature scales for creating different image representations, thereby increasing the similarity of the multi-modal images and thus co-registration accuracy.

Though there are a number of extant texture features (e.g. Haralick, Laws, Gabor), there also exists substantial redundancy between these features. The use of dimensional reduction (DR) methods to project high dimensional features into lower dimensional spaces allows for the construction of new transformed representations that can robustly drive multi-modal co-registration. Independent component analysis (ICA) is a method for decomposing mixture data into lower dimensional orthogonal features, with the most salient information obtained by minimizing mutual information (MI) and maximizing non-Gaussianity within the data. Li et al. showed that using ICA could reduce redundancy between multi-channel image texture features, in turn improving the co-registration result. ICA does not, however, implicitly rank the order of the independent components (ICs) according to their importance.

Spectral embedding (SE) is a nonlinear DR method for projecting data into a low-dimensional manifold from a high-dimensional space. This method takes the dissimilarity between two images as a weighted affinity matrix and applies eigenvalue decomposition to obtain the corresponding eigenvectors and eigenvalues. The approach orders the components that best preserve the structure of the high-dimensional manifold by selecting the eigenvectors corresponding to the minimum eigenvalues as the top components. In addition, applying SE onto the textural ICs extracted at each scale can help preserve the nonlinear relationships between the target and template images and hence potentially facilitate more accurate co-registration.

MI is a popular similarity measure for multi-modal image co-registration where the goal is to try to maximize the information shared between two images. MI mitigates the sensitivity of intensity-difference-based similarity measures on different image modalities by measuring intensity distribution of the images to be co-registered. Unfortunately, in spite of these advantages, MI is inadequate at robustly registering multi-modal images with substantially different appearance characteristics. For example, the appearance of tissue and anatomical structures on prostate MRI and corresponding ex vivo histopathology are vastly different. (Figure 1) The shape of the excised prostatectomy specimen tends to substantially change during the process of tissue fixation, slicing and sectioning.
The tissue preparation procedure has also been found to cause tissue loss and deformation. Similarly, the shape of in vivo MRI can undergo substantial deformation under pressure from surrounding organs such as the bladder. Furthermore, during MRI acquisition, the presence of an endorectal coil could also induce deformations in the natural shape of the prostate. Hence the goal of this work is to identify image representations which maximize the similarity between the fixed (MRI) and moving (histopathology) images. However, there is a need for higher dimensional similarity functions to be able to combine multiple different image representations to adequately co-register diverse looking images. To meet this need, we employ the $\alpha$-MI, which shows great performance in high dimensional registration as the similarity measure.

**Previous Work and Brief Overview of Approach**

As previously mentioned, there has been substantial interest in identifying alternative image representations for multi-modal co-registration. Chappelow et al. presented a texture-feature-based registration method to improve multi-modal registration performance called multi-attribute combined mutual information (MACMI). The approach used a set of multiple image texture features to complement image intensity information including the first and second order statistical and gradient features. MACMI yielded a noticeable improvement in alignment accuracy between images of different modalities, evaluations having been performed on both synthetic and clinical studies. Multi-modal co-registration in high dimensional spaces is difficult to drive using traditional MI approximated by the 'histogram-based plug-in' estimation. Thus, MACMI typically could not accommodate more than 2D or 3D spaces for co-registration. Instead of depending on density estimation, Neemuchwala et al. computed the length of entropic graphs, such as the $k$-Nearest Neighbor graph ($k$ NNG) and the minimal spanning tree (MST), to estimate graph entropic similarity measures. The graph entropic similarity measures, such as $\alpha$-MI, can potentially eliminate the low dimensionality constraint of 'histogram-based plug-in' estimation during calculation of MI. Staring et al. applied entropy graphs to address the deformable registration of cervical MRI and solved the high dimensional registration problem by deriving the analytic derivative of $\alpha$-MI with respect to the transformation parameters.

Furthermore, ICA has been employed to eliminate redundant representations from within a larger set of extracted textural features. However, the disadvantage of ICA is its unordered output ICs, hence it is impossible to fix the alternative representations for the fixed and moving images and thus the registration performance will most likely not be robust in spite of the repeated application of ICA. Hence, we employ spectral embedding onto the ICA representations. This enables generation of a series of ordered spectral vectors which are arranged in order of the amount of variance captured in the ICA component space. Thus, by having a ranked set of vectors, we can employ the top spectral embedding vectors to drive the co-registration. In addition, ICA is a linear DR method. For the MRI and ex vivo pathology specimens that we are attempting to co-register, a linear embedding representation may not adequately capture the non-linear deformations induced in both the imaging and the pathology data during the acquisition process. Spectral embedding is a nonlinear DR method and hence the
resulting vectors could potentially better capture the non-linear deformations induced in the imaging and pathology images.

In this paper, we present a new co-registration method, MSERg, that employs textural features, ICA, spectral embedding to identify the most representative feature scales and then combines these new multi-scale image representations via $\alpha$-MI to facilitate more accurate multi-modal co-registration. Specifically, our approach explores the influence of scale length of extracted texture features on multi-modal co-registration. Figure 2 provides an illustrative rendering of the overall workflow comprising our method.

Co-registration performance was compared with intensity-based registration, a texture-feature-based registration, named as multi-attribute combined mutual information (MACMI) registration 18, scale-invariant feature transform (SIFT) based registration, named as SIFT flow 19, and single-scale Spectral embedding representation registration (SSERg). SIFT flow is a registration algorithm developed by Liu et al. 19 and it employs the features extracted by SIFT descriptors to characterize the correspondence between the fixed and moving images. SIFT flow can be used to register images of great textural and appearance difference 19. Our experiment comprised of clinical and synthetic datasets. In the clinical experiment, we applied the Dice similarity coefficient (DSC) to evaluate the alignment of prostate capsule boundaries between the histopathology images and corresponding MRI which segmented by radiologists. For local registration accuracy evaluation, DSC was employed to evaluate the registration accuracy by comparing the registration results against the tumor annotations mapped from histopathology images via manual registration. In addition, the root mean squared distance (RMSD) was used.
to measure the distance between the corresponding landmarks between the histopathology images and MRI in millimeters. In the synthetic experiments, we firstly applied synthetic non-linear B-splines deformations on the proton density (PD) images and then registered the corresponding T1 weighted and T2 weighted images in order to recover the original applied deformation. We then compared the recovered deformation against the induced deformation to evaluate the performance of the co-registration scheme. The mean deformation difference (MDD) was employed to quantify the recovered deformation errors compared to the induced deformation.

**Results**

**Similarity improvement.** Figure 3 panel B shows a box plot of NMI distribution among 45 pairs of MRI and histology images with original intensity representation and the SE representations with scale \( \kappa \in \{3, 5, 7, 9, 11, 13, 15, 17\} \). Because each individual scale SE representation consists of 3 SE vectors, we use the average NMI value of these 3 corresponding SE vectors to represent the similarity level between SE representations of MRI and histology images. All the SE representations used in this experiment show greater NMI values than the intensity signal with moderate variance among different scales. The result indicates that SE representations could convert images of different modalities, MRI and histology images, to show more substantial similarity compared to the original intensity representation. In addition, SE representations of different scales extract different textural information. In Figure 3 panel A, (a) and (e) are the intensity representations of the moving and fixed images. (b–d) are histopathology SE representations and (f–h) are the corresponding MRI SE representations at scales \( \kappa \in \{3, 11, 17\} \), respectively. The SE representation at scale \( \kappa = 17 \) shows more similar appearance between the two modalities compared to other image representations. The influence of length scales on morphologic cues within the images is clearly illustrated. For instance, the region corresponding to the urethra is de-emphasized in the representations and at the length scales shown in (b) and (g). On the other hand, (c), (d), (f), and (h) highlight and emphasize the same regions. Panel C demonstrates the intensity representations of brain PD, T1 weighted and T2 weighted MRI in column (a) and SE representations at scale \( \kappa \in \{3, 9, 17\} \) respectively in columns (b)-(d). Similar to the clinical data shown in panel A, SE representations at different scales extract divergent morphology and texture features. Synthetic data SE representations in panel C column (c) illustrate underlying similarities across different modalities compared respectively to columns (a), (b) and (d). These results reinforce our approach, that different morphological structures are emphasized or suppressed across different length scales and

**Evaluation methods**

| Tumor lesion DSC | Intensity-based Registration | MACMI | SEFormer G | SIFT |
|------------------|-------------------------------|-------|------------|------|
| 0.59 ± 0.17      | 0.52 ± 0.18                   | 0.66 ± 0.10 | 0.54 ± 0.16 |
| Capsule boundary DSC | 0.93 ± 0.02               | 0.95 ± 0.03 | 0.96 ± 0.01 | 0.84 ± 0.04 |
| RSMD             | 3.97 ± 0.70                  | 3.64 ± 1.02 | 2.96 ± 0.76 | 3.89 ± 1.17 |
| MDD (T1 to PD)   | 33.78 ± 17.80                | 2.53 ± 1.05 | 1.45 ± 0.43 | 3.51 ± 0.73 |
| MDD (T2 to PD)   | 41.89 ± 16.19                | 1.82 ± 1.00 | 0.50 ± 0.28 | 1.36 ± 0.62 |

Table 1. Quantitative evaluation of representation-based registration on the clinical and synthetic testing sets. (N = 25 for both datasets) The table shows the mean and the standard derivation of tumor DSC, prostate boundary DSC and landmark RMSD for clinical dataset and MDD for synthetic dataset. Note that the greater the DSC, the lower the RMSD and MDD indicate the more accurate result.
only a multi-scale representation can optimally help determine image similarity in order to aid in multimodal co-registration.

**Evaluation of MSERg with other representation-based registration methods.** MSERg combined the scale $\kappa \in \{5, 7, 17\}$ for clinical data and the scale $\kappa \in \{3, 5, 7\}$ for synthetic data. These scale combinations were selected based on the evaluation of single-scale representation based registration performance on the independent learning set. For clinical data, the registration of the scale $\kappa \in \{3, 7, 17\}$ show averagely better tumor annotation alignment than other scales. For synthetic data, MMD values between the induced deformation and recovered deformation of the scale $\kappa \in \{3, 5, 7\}$ are less than other scales.

**MSERg and Intensity-based Registration.** For clinical data, MSERg illustrates a better across the board registration performance with less variance in the testing set compared with intensity-based registration. (Table 1) Furthermore, MSERg shows statistically significant performance improvement on all the three registration evaluation measurements. (Figure 4) Additionally, Figure 6 illustrates a clinical registration case showing intensity-based, MACMI, SIFT flow and MSERg registration results qualitatively. MSERg outperforms the other methods by showing the most accurate tumor ROI annotation, landmarks and capsule boundary alignment.

For synthetic data, the registration performance is evaluated via MDD. Table 1 and Fig. 5 demonstrate the quantitative performance evaluation among the four registration methods. MSERg has the least difference between the results and the induced deformation in both multimodal registration cases while intensity-based registration has the greatest difference.
MSERg and Texture-feature-based Registration. MACMI, a texture-feature-based-registration, although employs texture features like MSERg, the use of scale selection, ICA and SE appears confers a competitive advantage to MSERg. Figures 4 and 5 show the registration results with statistical significance analysis for clinical and synthetic dataset respectively. MSERg does not significantly outperform MACMI at a 95% confidence limit via landmark RMSD evaluation in clinical dataset. However, MSERg has significantly better boundary alignment and tumor annotation alignment accuracy.

MSERg and SIFT flow. SIFT is a descriptor to characterize image local intensity gradient information 20. SIFT flow adopts SIFT descriptors that aligns images of very different texture and appearances based on local gradient information 19. However, in case of in vivo T2 weighted MRI and ex vivo histology image registration, MSERg shows more accurate and robust results across 25 cases in the testing set. Especially, SIFT flow results have a poor boundary alignment accuracy compared with the other three registration methods. (Figures 4(a) and 6(j)) In addition, the MMD values of SIFT flow method in synthetic experiments are greater than MMD value of MSERg according to Figure 5. Thus, MSERg outperformed SIFT flow registration consistently on both clinical and synthetic datasets.

MSERg and SSERg. Figure 7 illustrates a frequency plot showing in how many instances either the SSERg or MSERg registration yielded the best result among clinical (N = 25) and synthetic (N = 25) testing sets. Although MSERg cannot outperform SSERg with every individual scale for every given experiment or dataset, overall MSERg dominates as the best performing registration scheme the majority of the time. The results clearly suggest that integrating multiple length scales offers an advantage in terms of multi-modal co-registration, the majority of the time.

Furthermore, the same scale combination kept in both T1 to PD and T2 to PD registrations and MSERg more frequently shows the best registration performance in both multimodal registration datasets. However, the results in Figure 7 also suggest that there is little consistency in performance of a specific scale across different experiments. In other words, there is no “magic scale”. For example, though scale $\kappa = 3$ is of frequency 0.32 as the best method in co-registering T1 to PD synthetic brain scans, scale $\kappa = 3$ does not feature even once when registering 25 pairs of T2 and PD synthetic brain scans. Thus, compared with SSERg, MSERg has more accurate and robust registration performance across different datasets.

Discussion
Co-registration of prostate in vivo MRI and ex vivo histopathology images is important in localizing prostate cancer extension on MRI and could help facilitate the development of radiomic machine based classifiers for characterization of prostate cancer on MRI 11,22. However, both the radiology and histopathology images undergo size and shape deformation during imaging and subsequent surgical resection and slice preparation, making automatic co-registration difficult. MSERg is an automatic registration method involving textural analysis scale selection, independent component analysis and spectral embedding to align the in vivo MRI and ex vivo histopathology images and thus map the ground truth tumor extension onto MRI.
The difference, however, between MSERg and the approaches of Li et al. of multimodal images can reduce dissimilarity between multimodal imagery and help facilitate co-registration within each monomodal image. The authors did not explicitly attempt to evaluate possible correlation and hence identify the optimal feature attributes in order to increase the multimodal image similarity and voxel uniqueness. MRI scans. Their approach requires high quality images and landmark points to constrain the deformation field. Features to fuse texture information from different image channels to drive multimodal co-registration of brain images.

However, we do acknowledge that one weakness of this study, albeit a direction of future work, is the intelligent scale selection. Currently we used a fairly naïve scale selection approach for constructing MSERg. The approach essentially involves selecting scales based on their average single-scale registration performance. Over the last decade there has been work on co-registration methods for pre-operative in vivo imaging and ex vivo histopathology images of the prostate. There are two general classes of methods to co-register in vivo MRI and ex vivo histopathology images of the prostate: 3D volume to volume registration and 2D slice to slice registration. 3D volume to volume registration requires for accurate 3D histology volume reconstruction. However, the 3D histology volume reconstruction is crippled by limited number of histology slices typically obtained during clinical workup of the surgical specimens. Consequently, there has been several approaches that have attempted 2D slice to slice registration. The limitation of the 2D slice to slice registration is the requirement for the slice correspondence to first be determined between the in vivo MRI and ex vivo histopathology images. Xiao et al. developed an iterative group-wise comparison methods to identify the correspondence between ex vivo histology and pre-operative in vivo MRI slices. However, the accuracy of the correspondence estimation tends to depend on the manual segmentation of the prostate on the MRI sections, which in turn depends on the radiologist who annotates the tumor. Another recent attempt at histology-MRI co-registration involves the construction of 3D printed molds containing spatial landmarks to relate the histology slices with in vivo MRI slices. While this approach could help address the constraints of establishing slice-to-slice correspondence, significant discrepancy might exist in the sagittal view between the 3D molds and MRI volume and it may not be possible to include this approach within a busy clinical protocol.

Other approaches for co-registering MRI and pathology images have involved either some manipulation of the pathology specimens (e.g. inserting carbon rods) or ex vivo imaging of the surgical specimen. The closest work to our approach is that of Li et al. Li et al. applied ICA on multi-scale and multi-oriented Gabor features to fuse texture information from different image channels to drive multimodal co-registration of brain MRI scans. Their approach requires high quality images and landmark points to constrain the deformation field. However, the prostate pathology and MRI scans such as the ones employed in our study may have various artifacts, making landmark selection subjective and user dependent. Ou et al. introduced a general-purpose multi-feature registration approach. They too employed Gabor filters and attempted to use feature selection approaches to identify the optimal feature attributes in order to increase the multimodal image similarity and voxel uniqueness within each monomodal image. The authors did not explicitly attempt to evaluate possible correlation and hence redundancy between selected filters and limited the textural representations to only the Gabor filter family.

The underlying rationale behind MSERg, just like in Li et al. and Ou et al. is that alternative representations of multimodal images can reduce dissimilarity between multimodal imagery and help facilitate co-registration. The difference, however, between MSERg and the approaches of Li et al. and Ou et al. is that MSERg involves the use of spectral embedding on independent components derived from multiple classes of texture features (Haralick and Gabor) to further distill and rank the textural information to construct alternative representations of the original images. Figure 3 illustrates that spectral embedding representations appear to be more capable of improving the similarity between the fixed and moving images compared to the original intensity representation. Furthermore, MSERg aims to use the spectral embedding representations of different size scales to highlight different types of details that may be useful in identifying similarities between dissimilar images and thus improve registration performance. For example, small scale texture features from MRI and histopathology images may reflect capillary vessel infiltration, or small scale tissue differentiation, while large scale texture feature filters may detect larger blood vessels, bone, or boundaries between organs. MSERg specifically aims to explore the influence of length scale of textural representation, and thus the registration performance.

It has been shown that MSERg can improve the radiology and histopathology image co-registration performance. However, we do acknowledge that one weakness of this study, albeit a direction of future work, is the intelligent scale selection. Currently we used a fairly naïve scale selection approach for constructing MSERg. The approach essentially involves selecting scales based on their average single-scale registration performance on the independent learning set. However, this naïve scale selection method still gave us very promising and encouraging results. Clearly though more intelligent scale selection approaches can be employed. Future work could potentially employ optimization techniques for more efficient and intelligent scale combination selection.

**Figure 7.** Illustration of a frequency plot showing in how many instances either SSERg or MSERg registration yielded the best registration result. We compared 8 different SSERg combinations along with MSERg on the same 25 clinical and synthetic testing cases. (a) Shows the frequency of each registration method having the greatest Tumor DSC within the clinical testing set. (b) and (c) illustrate the frequency of each registration method with the least mean deformation difference (MDD), in case of T1 to PD and T2 to PD registration respectively. Compared to SSERg, MSERg has more accurate and robust registration performance across different datasets.
Another limitation of this work was that a precise and detailed sensitivity analysis of multiple different parameters involved in MSERg was not performed. For instance, we did not evaluate the potential variations in performance as a function of the number of features for ICA, the number of ICs for SE and the number of SE vectors for each SE representation. This is another direction for future work. Finally, though the learning and testing sets used in this study were from two different institutions, more validation experiments need to be performed on multiple datasets from different institutions in order to more thoroughly evaluate the robustness of MSERg.

**Data Description and Experimental Design**

**Dataset Description.** This study consists of both a clinical as well as a synthetic dataset. In the clinical dataset, we considered 45 pairs of *in vivo* prostate histology specimens and corresponding 3 Tesla T2-w axial MRI from a total of 19 patients. Histology specimens were acquired after radical prostatectomy and sliced at 3 mm intervals. Each slice was stained with Hematoxylin and Eosin. The histology images were originally collected in quadrants and stitched into pseudo whole mount slices using Histostitcher® software. All data was analyzed retrospectively, after de-identification of all patient sensitive information. All experimental protocols were approved under the IRB protocol #02–13–42C with the University Hospitals of Cleveland Institutional Review Board, and all experiments were carried out in accordance with approved guidelines. Under this IRB, we were allowed to obtain de-identified images from St. Vincent's Hospital and University of Pennsylvania, and material transfer agreements were signed and agreed upon between Case Western Reserve University and University of Pennsylvania and St. Vincent's Hospital. Correspondences between histology and MRI and cancer region annotations on the histology image and MRI were made by a pathologist and a radiologist working together in unison. Here, we treated the MRI as the fixed image and corresponding histology image as the moving image.

The synthetic dataset contains 50 corresponding slices of 181 × 217 PD, T1 weighted and T2 weighted brain MRI from the MNI BrainWeb simulated brain database. The slice thickness of the dataset was 1 mm and intensity non-uniformity was 0%. We applied synthetic non-linear B-splines deformations on the PD images and then registered the corresponding T1 weighted and T2 weighted images in order to recover the original applied deformation. We then compared the recovered deformation against the induced deformation to evaluate performance of the co-registration scheme.

**Pre-processing of MRI and Pathology Data.** The prostate regions of T2-w MRI slices were manually segmented using 3D Slicer. The RGB digital images were converted to grayscale using MATLAB rgb2gray function, padded and down sampled to about 10 μm pixel resolution to match the resolution of the reference MRI. Each pair of pre-processed MRI and histology images were spatially concatenated to make sure that the same ICs were extracted on both the MRI and histology images. Texture feature extraction, ICA and SE were applied on the concatenated images.

**Experimental Design.** *Single-scale Representation Registration.* In our study, we constructed new SE representations for 8 different scale κ to concentrate texture information with 8 different sizes of neighborhood. The normalized mutual information (NMI) was treated as an evaluation measure to evaluate the improvement in similarity of SE representations compared against the original MRI and the signal intensity from the histology slices alone. This single-scale spectral embedding representation based registration employed single-scale SE representations for the moving and fixed images and was performed on 20 pairs of MRI and histology images and 25 pairs of synthetic images in the learning set and then the registration results were evaluated in terms of DSC for clinical data and MDD for synthetic data at each κ ∈ [3, 5, 7, 9, 11, 13, 15, 17]. The top 3 performing scale representations were selected for MSERg based on the average registration performance in the learning set.

**Multi-scale representation.** The top 3 individual scale representations identified in the learning set were combined in a high dimensional registration space (Figure 2). MSERg employs α-MI as the similarity measure to conduct high dimensional registration on both the clinical and synthetic testing sets. The goal of this experiment was to evaluate whether combining the top performing individual scale representations can further improve multi-modal registration performance. In this work we compared MSERg with four other registration-based registration strategies, intensity-based registration, texture-feature-based registration, SIFT flow registration and single-scale representation based registration.

**Intensity-based registration, texture-feature-based registration and SIFT flow registration.** Here, ‘intensity-based registration’ refers to registration using the original intensity images as the fixed and moving images with MI estimated on the basis of a histogram as the similarity measure. The registration parameter selection and optimization were done during the learning process and on the training step. The set of parameters that resulted in the best registration result were locked down for the evaluation on the hold out testing set. For the texture-feature-based registration, we adopt MACMI method to selected the texture feature representations. MACMI selects the texture features that maximize the combined mutual information (CMI). We extracted an ensemble of five features with optimal CMI based on the same gradient, first and second order statistical features used in the work of Chappelow et al. Following feature selection, free form deformed registration was applied with α-MI as the similarity measure to co-register the multi-modal images in the testing set. In addition, we have validated with SIFT flow registration as proposed in the work of Liu et al.

**Ground Truth Registration.** In order to quantitatively evaluate the performance of the different co-registration methods in the clinical dataset, we need a ground truth definition of what constitutes a near optimal registration. This can then be used as the framework for comparing all other registration results. Towards this end, a control-point-based manual registration scheme was employed for co-registering all 45 pairs of histology-MRI
sections. The corresponding control points were carefully selected by a pathologist and a radiologist on ex vivo histology images and in vivo MRI. The landmarks used for each registration pair varied between 6 to 10. The registration results were then visually assessed by the radiologist and pathologist. If they were found to be unsatisfactory, the registration was re-done. Once the co-registrations were found to be satisfactory, they were designated as the "ground truth" registration for the particular MRI-pathology pair.

Performance Evaluation Measures. Registration results were assessed via DSC⁴ and RMSD. DSC is defined as follows,

\[ DSC(X_0, X_1) = \frac{2|A_{X_0} \cap A_{X_1}|}{|A_{X_0}| + |A_{X_1}|}, \]

where \( A_{X_0} \) denotes the region of interest (ROI) previously manually delineated on the fixed images. Here ROI refers to the capsule segmentation in global accuracy evaluation and the tumor annotated regions on the fixed images in local accuracy evaluation. \( X_0 \) and \( X_1 \) refer to the fixed and moving images respectively. The tumor annotations on the fixed image are obtained via manual registration between the histopathology images and MRI. \( A_{X_1} \) refers to corresponding annotations on the moving images.

RMSD is defined as follows,

\[ RMSD(X_0, X_1) = \sqrt{\frac{1}{K} \sum_{k=1}^{K} \|X_0(\beta_k) - X_1(\beta_k)\|^2}, \]

where \( \beta_k \) denotes the pixel of \( k \)th landmark in \( X_0 \) and \( X_1 \) and \( \| \cdot \| \) represents the \( L_2 \) norm. RMSD measures the distance between the anatomical landmarks such as the urethra, and nodules annotated by the pathologist and radiologist.

The MDD refers to

\[ MDD(D_0, D_1) = \frac{\sum_{k=1}^{C} (D_0 - D_1)}{C}, \]

where \( D_0 \) is the ground true deformation field and \( D_1 \) is the result deformation field. \( C \) is the total number of deformation vectors in the deformation field. MDD measures the difference between the registration result and the ground truth.

Implementation Details. SE representations. To construct SE representations for different length scales, firstly, Gabor and Haralick features were extracted for \( \kappa \in \{3, 5, 7, 9, 11, 13, 17\} \). Then ICA was employed on the Gabor and Haralick features across different \( \kappa \) using the JADE-ICA algorithm. Finally, the spectral embedding algorithm was used to get the top 3 SE vectors for each individual scale representation.

Identifying optimal scales. In the learning set, single-scale SERg co-registration was implemented within each individual scale SE space. \( \alpha \)-MI was applied as the high dimensional similarity measure with \( k \)-NNG as the entropic graph. The deformable registration was realized via the Elastix toolbox⁵ with B-spline interpolation. We set \( \alpha = 0.99, k = 20 \) nearest neighbors with 3 resolution levels (resolution scales \( \eta = 4.0, 2.0, 1.0 \)) employed with 100 iterations for each resolution level. The top 3 representations with the greatest DSC were selected as the optimal scale representations for MSERg.

Comparing MSERg against other representation-based registration methods. All the registration methods adopted the same standard gradient descent optimization method via the Elastix toolbox⁵. MSERg was compared against intensity-based registration and texture-feature-based registration. For intensity-based registration, 3 resolution levels (resolution scales \( \eta = 4.0, 2.0, 1.0 \)) were employed with 1000 iterations for each resolution level. The texture-feature-based registration also were made to use the same multi-resolution registration strategy with 100 iterations for each resolution level. MSERg employed a combination of 3 scale SE representations (\( \kappa \in \{5, 7, 17\} \)), with \( \alpha = 0.99, k = 20 \) for \( \alpha \)-MI registration in Elastix and 100 iterations per resolution level (resolution scales \( \eta = 4.0, 2.0, 1.0 \)). SIFT flow was implemented via the Image Alignment Toolbox (IAT), which is a Matlab toolbox for image alignment and registration⁶. The manual registration to obtain the ground truth for the different schemes was implemented via an in-house software tool using thin-plate splines⁷ method.

Computational efficiency. For a pair of MRI and histology images with 320 × 320 resolution, the spectral embedding representation construction for all 8 scales takes around 10 minutes and the representation construction for MACMJI and SIFT takes 5 minutes and 7 seconds respectively. In the registration step, each single-scale spectral embedding representation registration requires around 1 minute to align a pair of MRI and histology images. Thus, it takes 1 × 8 × 20 = 160 minutes for all 8 single-scale representations in the learning set consisting of 20 cases. This computational burden can be greatly reduced via parallel computing because all the single-scale representation registrations are independent of each other. In the testing set, MSERg, MACMJI and intensity-based registration take around 2.5, 3.5 minutes and 0.5 minute respectively. All three registration methods were
implemented in the Elastix toolbox on a computer with a Windows 10 operation system based off a 3.40 GHz processor with 16.0 GB RAM. SIFT registration was implemented via SIFT flow algorithm taking about 1.5 minutes for each case on the same computer, using the same software and OS configuration. Table 2 summarizes the computational efficiency for all registration methods used in this paper.

### Table 2. Computational efficiency including the image representation calculation and registration time consumption. Although MSERg has a higher computational cost during the learning step, this burden can be greatly reduced via parallel computing because all the single-scale representation registrations are essentially independent of each other, with no dependencies.

| Registration methods | Intensity-based | MACMI | MSERg | SIFT |
|----------------------|----------------|-------|-------|------|
| Representation computation | NA | 5 min | 10 min | 7 sec |
| Registration | 0.5 min | 3.5 min | 160 min for learning; 2.5 min for testing | 1.5 min |

**References**

1. Chappelow, J., Tomaszewski, J. E., Feldman, M., Shih, N. & Madabhushi, A. Histostitcher ©: An interactive program for accurate and rapid reconstruction of digitized whole histological sections from tissue fragments. *Computerized Medical Imaging and Graphics* 35, 557–567 (2011).

2. Pluim, J. P., Maintz, J. A. & Viergever, M. A. Image registration by maximization of combined mutual information and gradient information. In *Medical Image Computing and Computer-Assisted Intervention–MICCAI* 2000, 452–461 (Springer, 2000).

3. Rueckert, D., Clarkston, M., Hill, D. & Hawkes, D. J. Non-rigid registration using higher-order mutual information. In *Medical Imaging* 2000, 438–447 (International Society for Optics and Photonics, 2000).

4. Studholme, C., Hill, D. & Hawkes, D. Incorporating connected region labelling into automated image registration using mutual information. In *Mathematical Methods in Biomedical Image Analysis*, 1996., Proceedings of the Workshop on, 23–31 (IEEE, 1996).

5. Daugman, J. G. Complete discrete 2-d gabor transforms by neural networks for image analysis and compression. *Acoustics, Speech and Signal Processing, IEEE Transactions on* 36, 1169–1179 (1988).

6. Li, Y. & Verma, R. Multichannel image registration by feature-based information fusion. *Medical Imaging, IEEE Transactions on* 30, 707–720 (2011).

7. Ou, Y., Sotiras, A., Paragios, N. & Davatzikos, C. Dramms: Deformable registration via attribute matching and mutual-saliency weighting. *Medical image analysis* 15, 622–639 (2011).

8. Chappelow, J., Madabhushi, A., Rosen, M., Tomaszewski, J. & Feldman, M. A combined feature ensemble based mutual information scheme for robust inter-modal, inter-protocol image registration. In *Biomedical Imaging: From Nano to Macro*, 2007. ISBI 2007. 4th IEEE International Symposium on, 644–647 (IEEE, 2007).

9. Chappelow, J. et al. Elastic registration of multimodal prostate mri and histology via multiattribute combined mutual information. *Medical Physics* 38, 2005–2018 (2011).

10. Hwuang, E. et al. Spectral embedding-based registration (serg) for multimodal fusion of prostate histology and mri. In *SPIE Medical Imaging*, 90343P–90343P (International Society for Optics and Photonics, 2014).

11. Miyamoto, E. & Merryman, T. Fast calculation of haralick texture features. *Human computer interaction institute, Carnegie Mellon University, Pittsburgh, USA. Japanese restaurant office* (2005).

12. Laws, K. I. Textured image segmentation. *Tech. Rep., DITIC Document* (1980).

13. Hyvarinen, A. & Oja, E. Independent component analysis: algorithms and applications. *Neural networks* 13, 411–430 (2000).

14. Pluim, J. P., Maintz, J. A. & Viergever, M. et al. Mutual-information-based registration of medical images: a survey. *Medical Imaging, IEEE Transactions on* 22, 986–1004 (2003).

15. Staring, M. et al. Registration of cervical mri using multifeature mutual information. *Medical Imaging, IEEE Transactions on* 28, 1412–1421 (2009).

16. Viola, P. & Wells, W. M. III Alignment by maximization of mutual information. *International journal of computer vision* 24, 137–154 (1997).

17. Neemuchwala, H., Hero, A., Zabuawala, S. & Carson, P. Image registration methods in high-dimensional space. *International Journal of Imaging Systems and Technology* 16, 130–145 (2006).

18. Chappelow, J. & Madabhushi, A. Multi-attribute combined mutual information (macmi): An image registration framework for leveraging multiple data channels. In *Biomedical Imaging: From Nano to Macro, 2010 IEEE International Symposium on*, 376–379 (IEEE, 2010).

19. Liu, C., Yuen, J. & Torralba, A. Sift flow: Dense correspondence across scenes and its applications. *IEEE transactions on pattern analysis and machine intelligence* 33, 978–994 (2011).

20. Lowe, D. G. Object recognition from local scale-invariant features. In *Computer vision, 1999. The proceedings of the seventh IEEE international conference on*, vol. 2, 1150–1157 (Ieee, 1999).

21. Hoeks, C. M. et al. Prostate cancer: multiparametric mr imaging for detection, localization, and staging. *Radiology* 261, 46–66 (2011).

22. Viswanath, S. et al. Integrating structural and functional imaging for computer assisted detection of prostate cancer on multi-protocol in vivo 3 tesla mri. In *SPIE medical imaging*, 72603I–72603I (International Society for Optics and Photonics, 2009).

23. Orczyk, C. et al. Imaging of prostate cancer: A platform for 3d co-registration of in-vivo mri ex-vivo mri and pathology. In *SPIE Medical Imaging*, 83162M–83162M (International Society for Optics and Photonics, 2012).

24. Park, H. et al. Registration methodology for histological sections and in vivo imaging of human prostate. *Academic radiology* 15, 1027–1039 (2008).

25. Xiao, G. et al. Determining histology-mri slice correspondences for defining mri-based disease signatures of prostate cancer. *Computerized Medical Imaging and Graphics* 35, 568–578 (2011).

26. Priester, A. et al. A system for evaluating magnetic resonance imaging of prostate cancer using patient-specific 3d printed molds. *American journal of clinical and experimental urology* 2, 127 (2014).

27. Groenendaal, G. et al. Validation of functional imaging with pathology for tumor delineation in the prostate. *Radiotherapy and Oncology* 94, 145–150 (2010).

28. Chappelow, J., Madabhushi, A., Rosen, M., Tomaszewski, J. & Feldman, M. Multimodal image registration of ex vivo 4 tesla mri with whole mount histology for prostate cancer detection. In *Medical Imaging*, 65121S–65121S (International Society for Optics and Photonics, 2007).*
29. Ohnishi, T. et al. Deformable image registration between pathological images and MR image via an optical macro image. *Pathology-Research and Practice* **212**, 927–936 (2016).

30. Cocosco, C. A., Kollokian, V., Kwan, R. K.-S., Pike, G. B. & Evans, A. C. Brainweb: Online interface to a 3d mri simulated brain database. *In NeuralImage* (Citeseer, 1997).

31. Pieper, S., Halle, M. & Kikinis, R. 3d slicer. *In Biomedical Imaging: Nano to Macro, 2004. IEEE International Symposium on*, 632–635 (IEEE, 2004).

32. Rueckert, D. et al. Nonrigid registration using free-form deformations: application to breast mr images. *IEEE transactions on medical imaging* **18**, 712–721 (1999).

33. Zou, K. H. et al. Statistical validation of image segmentation quality based on a spatial overlap index 1: Scientific reports. *Academic radiology* **11**, 178–189 (2004).

34. Klein, S. et al. Elastix: a toolbox for intensity-based medical image registration. *Medical Imaging, IEEE Transactions on* **29**, 196–205 (2010).

35. Image alignment toolbox (iat), a matlab toolbox for image alignment and registration. http://iatool.net/.

36. Bookstein, F. L. Principal warps: Thin-plate splines and the decomposition of deformations. *IEEE Transactions on Pattern Analysis & Machine Intelligence* **11**, 567–585 (1989).

**Acknowledgements**

Research reported in this publication was supported by the National Cancer Institute of the National Institutes of Health under award numbers R21CA179327-01, R21CA195152-01, U24CA199374-01 the National Institute of Diabetes and Digestive and Kidney Diseases under award number R01DK098503-02, the DOD Prostate Cancer Synergistic Idea Development Award (PC120857); the DOD Prostate Cancer Idea Development Award; the Case Comprehensive Cancer Center Pilot Grant the VelaSano Grant from the Cleveland clinic the Wallace H. Coulter Foundation Program in the Department of Biomedical Engineering at Case Western Reserve University the I-Corps@Ohio Program Case Urology Translational Research Training Program (CUTRTP) Hartwell Foundation. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

**Author Contributions**

L.L. performed all experiments, wrote the manuscript text, and generated the tables and figures. A.M. provided supervision of the work; S.P. and J.G. provide the annotations on the MRI and histopathology images; G.P. conducted the manual registration; S.V. and M.R. provide the algorithm suggestions to the experiments.

**Additional Information**

Supplementary information accompanies this paper at doi:10.1038/s41598-017-08969-w

**Competing Interests:** The authors declare that they have no competing interests.

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