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Spectral clustering to model deformations for fast multimodal prostate registration

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Abstract

This paper proposes a method to learn deformation parameters off-line for fast multimodal registration of ultrasound and magnetic resonance prostate images during ultrasound guided needle biopsy. The method is based on a learning phase where deformation models are built from the deformation parameters of a spline-based non-linear diffeomorphism between training ultrasound and magnetic resonance prostate images using spectral clustering. Deformation models comprising of the eigen-modes of each cluster in a Gaussian space are applied on a test magnetic resonance image to register with the test ultrasound prostate image. The deformation model with the least registration error is finally chosen as the optimal model for deformable registration. The rationale behind modeling deformations is to achieve fast multimodal registration of prostate images while maintaining registration accuracies which is otherwise computationally expensive. The method is validated for 25 patients each with a pair of corresponding magnetic resonance and ultrasound images in a leave-one-out validation framework. The average registration accuracies i.e. Dice similarity coefficient of 0.927 ± 0.025, 95% Hausdorff distance of 5.14 ± 3.67 mm and target registration error of 2.44 ± 1.17 mm are obtained by our method with a speed-up in computation time by 98% when compared to Mitra et al. [7].

1. Introduction

The appearance of malignant lesions in a Transrectal Ultrasound (TRUS) guided needle biopsy of the prostate is mostly hypoechoic and the accuracy of finding such lesions is typically 43% in sonography. Approximately 25% - 42% of cancer lesions can also be isoechic and the chance to diagnose hypoechoic malignant lesions from TRUS guided biopsy is ≤ 57% [3]. Magnetic Resonance (MR) images provide better soft-tissue contrasts, therefore, fusion of pre-biopsy MR images onto interventional TRUS images might increase the overall biopsy accuracy [9].

The prostate of the same patient may exhibit deformations between the TRUS and the MR images. The deformations are caused by the insertion of the endorectal probe during the MR acquisition, bowel or gas inside the rectum and displacement of patient position between the TRUS and MR imaging procedures. However, the deformation is mostly observed as flattening of the part of the prostate adjacent to the rectum. Therefore, in this paper we attempt to model such deformations of the prostate from a set of corresponding MR images co-registered with the TRUS images. The deformable registration of the training set of TRUS and MR images is done using the method of Mitra et al. [7]. However, instead of the geometric approach used in their work to establish point correspondences, we employ the shape-context based method of Belongie et al. [1] and Bhattacharyya distance [2] to set contour point correspondences across the TRUS and the MR prostate images.

The deformable registration in [7] is based on the minimization of the difference in segmented prostate regions where both the TRUS and MR regions are under the influence of a set of polynomial functions. The MR image transformation employs a thin-plate spline (TPS) interpolation. The combination of TPS based interpolation and the set of polynomial functions ensures a smooth diffeomorphic transformation of the MR image at the cost of increased computation time. However, the MR images need to be registered with the TRUS images in near real time during prostate biopsy. Therefore, to achieve fast registration we propose to model the TPS weight parameters obtained from the diffeomorphic registration of training TRUS-MR images and then apply the modeled parameters to register a new set of TRUS-MR images. A single deformation model in
Gaussian space derived from the principal eigen-modes of the deformation vectors as shown by [5] is not sufficient to model all the variations of prostate deformation. Therefore, we propose to cluster the deformation vectors into similar groups using spectral clustering approach. The principal eigen-modes of the deformation vectors of each of these clusters in a Gaussian space form a deformation model. The registration of a test set of TRUS-MR images involves recovering the affine parameters from the established point correspondences and the TPS weight parameters of each of the deformation model are consecutively applied. The model with the least registration error between the TRUS-MR images is chosen as the optimal deformation model.

The remaining paper is organized in the following manner. Section 2 describes the proposed method. Section 3 provides the results related to registration accuracies and the computation time, followed by the conclusions in Section 4.

2. The Proposed Method

The proposed method is based on the following components: 1) Point correspondences established on both the TRUS and MR images that are required for both the training and testing phases, 2) the non-linear diffeomorphic framework required for deformation of the training MR images, 3) spectral clustering of TPS weight vectors during the training and 4) linear estimation of deformation parameters applied on the test MR image. The following paragraphs provide descriptions of the afore-mentioned components.

Point Correspondences: In this work we use manually segmented prostate shapes for accurate evaluation of our method. However, in future we plan to use some automatic prostate segmentation methods in TRUS and MRI such as proposed in [6]. The segmented prostate contour points are uniformly sampled using fixed Euclidean distance. Let the number of uniformly sampled points now be represented as \( n \), then each sample point \( c_i \) may be represented by a shape descriptor that is a \( n - 1 \) length vector of log-polar relative distances to points \( c_j \), where \( i \neq j \). The shape descriptor is binned into a histogram that is uniform in log-polar space and this histogram is the shape-context representation of a contour point [1] i.e. \( c_i \) is represented by a histogram \( h_i(k, \theta) \) such that

\[
 h_i(k, \theta) = \# \{ c_j, i \neq j : (c_i - c_j) \in \text{bin}(k, \theta) \}, \quad (1)
\]

where \( k = \tan^{-1}\frac{x_{j2}-x_{i2}}{x_{i1}-x_{j1}} \) of the relative distance \( (c_i - c_j) \), where, \( c_i = (x_{i1},x_{i2}) \) and \( c_j = (x_{j1},x_{j2}) \). As suggested by Belongie et al. [1], a total of 5 bins are considered for \( k \) and 12 bins for \( \theta \) that ensures that the histogram is uniform in log-polar space.

In this work we choose the Bhattacharyya distance [2] between the shape-context histograms of two shapes to find the best point correspondence since it is fast to compute. Thus, to match a point \( c_i \) in a shape to a point \( c_j \) in another shape, the Bhattacharyya coefficients between the shape-context histogram of \( c_i \) and all \( c_j \) are computed and the \( c_j \) that maximizes the relation in Eq. (2) is chosen as the corresponding point.

\[
 \arg \max_{c_j} \sum_{k=1}^{5} \sum_{\theta=1}^{12} \sqrt{h_i(k,\theta)h_j(k,\theta)}, \quad (2)
\]

where, \( h_i(k,\theta) \) and \( h_j(k,\theta) \) are the normalized shape-context histograms of \( c_i \) and \( c_j \) respectively.

The smoothness of the transformation between the MR and TRUS images may be guaranteed if the prostate mask centroids are also considered in addition to the contour correspondences. The set of correspondences established will be referred as \( p_i^m \) and \( p_i^f \), where \( i = 1,2,\ldots,9 \) for the moving and fixed images respectively in the following sections. Fig. 1 shows the 8 contour correspondences and the centroid of the prostate overlaid on the TRUS and MR images.

Non-linear Diffeomorphism: Point correspondences established on a pair of training TRUS-MR images are used to align the moving MR image with the fixed TRUS image. Assuming that \( x = [x_1,x_2] \in \mathbb{R}^2 \) and \( y = [y_1,y_2] \in \mathbb{R}^2 \) are the moving MR and fixed TRUS images respectively, a system of nonlinear equations is constructed as proposed in [4, 7]:

\[
 \int_{I_y} \omega_k(y)dy = \int_{I_x} \omega_l(\varphi(x)) |J_\varphi(x)| dx, \quad (3)
\]

\( \varphi(x) = [\varphi_1(x), \varphi_2(x)] \) is the deformation field, \( |J_\varphi(x)| \) is the Jacobian determinant of the transformation at \( x \) of the moving image. Each nonlinear function \( \omega_l(\cdot), \) \( l = 1,\ldots,l \) generates one equation, yielding a system of \( l \) equations [4]. The transformation \( \varphi(x) \) is a TPS interpolation of the moving image guided by the set of points established on the same and is given by Eq. (4) where, \( a_{iv} \) are the 6 affine parameters, \( w_{iv} \) are...
the TPS weight parameters for the control points with
\( i = 1, \ldots, n, v = 1, 2, u = 1, 2, 3, U(r) = r^2 \log r^2 \) is
the radial-basis function and \( \| \cdot \| \) is the Euclidean norm.
\[
\varphi_v(x) = a_{v1} x_1 + a_{v2} x_2 + a_{v3} + \sum_{i=1}^{n} \, w_{iv} U(||p_i - x||),
\]
s.t. \( \sum_{i=1}^{n} w_{iv} = 0 \) and \( \sum_{i=1}^{n} P_i^T w_{iv} = 0 \quad (4) \)
The diffeomorphic framework also considers the correspon-
dence localization error across the moving and
fixed images and the regularization of the bending en-
ergy of the TPS as
\[
\frac{1}{n} \sum_{i=1}^{n} \frac{||p_i^t - \zeta(p_i^m)||^2}{\sigma_i^2} + \lambda E_{TPS} = 0, \quad (5)
\]
where \( \sigma_i^2 \) is the sum of the variances of both the mov-
ing and fixed point correspondences, \( E_{TPS} = \Delta^2\zeta \)
with \( \zeta(p_i^m) = p_i^t \). The adopted set of non-linear functions are the power functions \( \omega_i(x) = x_i^{\alpha_i} x_j^{\beta_j} \),
with \( (\alpha_i, \beta_j) \in \{(0,0), (0,1), (1,0), (1,1), \ldots, (5,5)\} \)
which provides sufficiently many equations to obtain
a least squares estimate of the 9 \times 2 TPS weights and
6 affine parameters. The solution of Eq. (3) with the
constraints in Eq. (4) and Eq. (5) is then obtained via
the Levenberg-Marquardt algorithm.

**Spectral clustering:** The deformation parameters
i.e. the TPS weight parameters obtained for the set
of training fixed and moving images are grouped into
similar deformation clusters by a spectral clustering
approach that determines the number of clusters auto-
matically. Cosine similarities of \( \mathcal{P} \) deformation vectors \( W = (w_1, w_2, \ldots, w_9) \) of the training set are used to
construct a \( \mathcal{P} \times \mathcal{P} \) similarity matrix \( W \). The objective
is to determine \( k \) disjoint clusters and the algorithm
may be defined in the following steps [8]:

1) Form the similarity matrix \( W \in \mathbb{R}^{P \times P} \).
2) Construct the normalized Laplacian \( L \) as \( D^{-1/2} W D^{-1/2} \) where \( D = \text{diag}(W) \).
3) Compute the first \( k \) eigenvectors of \( L \) to build the
matrix \( U \) by stacking the eigenvectors into columns.
4) Re-normalize the matrix \( U \) to make each row have
unit-length and apply K-means clustering to \( U \).
Similar deformation vectors are now grouped into
\( k \) disjoint clusters, where \( k \) is the number of largest
eigen-vectors comprising 88% of the total variations
such that each cluster consists of more than one
deforation vector.

**Linear estimation:** Geva et al. [5] showed an
off-line linear estimation of basis functions from a
deforation space. They performed a PCA of the coeffi-
cients of a bivariate B-splines transformation
to represent them by their principal eigen-modes.
Similarly, given a test moving image we may transform
it by the linear estimation of the TPS deformation
parameters i.e. \( w_{iv} = \sum_{s=1}^{N_s} a_s b_{i,s}^t \). \( N_s \) is the number
of principal axes on which the coefficients are projected
after PCA with \( a_s \) and \( b_{i,s}^t \) as the respective eigen-value
and the eigen-vector. Therefore, the transformation
\( \varphi(x) \) of Eq. (3) may be written as
\[
\varphi_v(x) = a_{v1} x_1 + a_{v2} x_2 + a_{v3} + \sum_{i=1}^{n} \sum_{s=1}^{N_s} a_s^t b_{i,s}^t U(||p_i^m - x||) \quad (6)
\]
The eigen-modes of \( k \) deformation clusters comprise
95% variation of the principal modes and its Gaus-
sian space of \(-2\sigma, -1\sigma, \ldots, +2\sigma\), with \( \sigma \) as the standard
deviation provide 5 deformation models for each
cluster. The affine parameters of the TPS transformation
are obtained by sum-of-squared differences minim-
ization of the point correspondences established on
the test moving and fixed images. Finally to obtain
the optimal transformation of the test moving image,
the registration error is computed as the percentage
of non-overlapping area in prostate foreground and the one
with least registration error is considered as the final
transformation.

### 3. Experiments and Results

The validation of our method is done using 25 pa-
tients axial mid-gland slices for both the TRUS and MR
images with an average size of 250 \times 200 pixels with
each pixel dimension being 0.2734 \times 0.2734 mm. A
leave-one-out approach is used where the deformation
models are formed from 24 datasets and are applied
to transform the remaining one. The registration accu-
racity is evaluated in terms of Dice similarity coefficient
(DSC) that measures the global overlap of the prostate
regions, 95% Hausdorff distance (HD) that measures
the contour accuracy and target registration error (TRE)
that measures the extent of overlap of the anatomical
targets in the transformed MR image and the TRUS
image. Table 1 shows the registration accuracies for the
method of Mitra et al. [7] with 20 patient datasets,
the proposed method with non-linear deformation on
improved point correspondences without deformation

<table>
<thead>
<tr>
<th>Methods</th>
<th>Mitra et al. [7]</th>
<th>DEF-NL</th>
<th>DEF-L</th>
</tr>
</thead>
<tbody>
<tr>
<td>DSC</td>
<td>0.98±0.004</td>
<td>0.97±0.010</td>
<td>0.92±0.25</td>
</tr>
<tr>
<td>HD</td>
<td>1.54±0.46</td>
<td>2.05±1.26</td>
<td>5.14±3.61</td>
</tr>
<tr>
<td>TRE</td>
<td>1.90±0.127</td>
<td>1.71±1.23</td>
<td>2.44±1.17</td>
</tr>
<tr>
<td>Time</td>
<td>320.79±76.01</td>
<td>106.34±32.45</td>
<td>4.99±3.52</td>
</tr>
</tbody>
</table>

---

Table 1. Registration accuracies and computation time for different methods. HD and TRE are in (mm) and time is in seconds.
learning abbreviated as DEF-NL and that with the deformation estimation (DEF-L) i.e. non-linear deformation applied to a set of training TRUS-MR images, spectral clustering to group deformations and thereby applying linearly estimated deformation parameters to transform a test moving MR for 25 datasets in a leave-one-out framework.

It is to be noted in Table. 1 that the method of Mitra et al. [7] using 37 correspondences needs more computation time than our proposed method with 9 point correspondences only. The registration accuracies of our method with 9 correspondences without the off-line deformation learning are comparable to [7]. The area and contour overlap accuracies in terms of DSC and HD for our method without the deformation learning approach are statistically significantly better with Student’s t-test \( p < 0.0001 \) and \( p < 0.001 \) respectively than that with the deformation learning. The TRE for our method without the deformation learning approach is also slightly better than that with the deformation learning with a statistical significance of \( p < 0.005 \). Nevertheless, the computation time of our method with deformation learning shows a statistically significant reduction with \( p < 0.0001 \) than the remaining methods. Fig. 2 shows the registration results for 2 patients, where it is observed that our method with and without deformation learning produce similar results. The accuracy of our method is qualitatively comparable with that of Xu et al. [9] that demonstrates a near real-time TRUS-MR prostate fusion method with an average registration error of \( 2.3 \pm 0.9 \) mm but requires \( 15 \) s for the registration process. Our method was implemented in MATLAB 2009(b) with 1.66GHz processor and 2GB memory. The method shows a significant speed-up of computation time when the off-line deformation learning approach is employed while maintaining a clinically significant average target registration error of \(< 3\) mm.

4. Conclusions

A method of deformable registration between TRUS and MR prostate images with prior learning of deformation parameters has been proposed. Spectral clustering has been used to group similar deformations from training TRUS-MR images and thereafter the eigen-modes of deformations for each deformation cluster in a Gaussian space have been used to deform a new MR image corresponding to the TRUS image. The method is fast and efficient to transform a moving image with good registration accuracy and may be used during prostate biopsy if programmed on GPU. The accuracy of resulting deformation may be further increased if more patient sets are used to learn the deformation parameters.

References