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Longitudinal, intermodality registration of quantitative breast PET and MRI data acquired before and during neoadjuvant chemotherapy: Preliminary results

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Purpose: The authors propose a method whereby serially acquired DCE-MRI, DW-MRI, and FDG-PET breast data sets can be spatially and temporally coregistered to enable the comparison of changes in parameter maps at the voxel level.

Methods: First, the authors aligned the PET and MR images at each time point rigidly and nonrigidly. To register the MR images longitudinally, the authors extended a nonrigid registration algorithm by including a tumor volume-preserving constraint in the cost function. After the PET images were aligned to the MR images at each time point, the authors then used the transformation obtained from the longitudinal registration of the MRI volumes to register the PET images longitudinally. The authors tested this approach on ten breast cancer patients by calculating a modified Dice similarity of tumor size between the PET and MR images as well as the bending energy and changes in the tumor volume after the application of the registration algorithm.

Results: The median of the modified Dice in the registered PET and DCE-MRI data was 0.92. For the longitudinal registration, the median tumor volume change was $-0.03\%$ for the constrained algorithm, compared to $-32.16\%$ for the unconstrained registration algorithms ($p = 8 \times 10^{-6}$). The medians of the bending energy were 0.0092 and 0.0001 for the unconstrained and constrained algorithms, respectively ($p = 2.84 \times 10^{-7}$).

Conclusions: The results indicate that the proposed method can accurately spatially align DCE-MRI, DW-MRI, and FDG-PET breast images acquired at different time points during therapy while preventing the tumor from being substantially distorted or compressed. © 2014 American Association of Physicists in Medicine. [http://dx.doi.org/10.1118/1.4870966]
Key words: registration, DCE-MRI, FDG-PET, breast cancer, prone PET, PET/MRI, longitudinal registration, treatment response

1. INTRODUCTION

In recent years there have been dramatic increases in both the quality and quantity of noninvasive imaging methods for assessing (and even predicting) the response of breast tumors to neoadjuvant therapy (NAT). In particular, dynamic contrast enhanced magnetic resonance imaging (DCE-MRI), diffusion weighted MRI (DW-MRI), and 18F-fluorodeoxyglucose positron emission tomography (FDG-PET) (Refs. 4 and 5) have matured to the point where each modality offers unique and, importantly, complementary information on several clinically relevant tumor characteristics. Typically, changes in these quantitative imaging parameters are summarized on a region of interest (ROI) basis which discards spatial information on tumor heterogeneity. However, there has been an increasing interest in the study of the tumor at the voxel level rather than the ROI level. Such analysis typically requires temporal and spatial registration of image datasets in order to measure voxelwise parameter changes in a meaningful way. In this effort, we present the first report which presents a method for rigorous registration of quantitative PET and MRI breast data acquired during NAT, which is a necessary step to allow for a more comprehensive analysis of tumor treatment response.

2. MATERIALS AND METHODS

2.A. Patient population and data acquisition

Data were acquired from ten patients with Stage II/III breast cancer enrolled in a IRB-approved clinical study prior to any treatment (t1), after one cycle of NAT (t2), and at the completion of NAT (t3). The data were not available at t3 for five of the patients. Table I provides the clinical characteristics of the patients included in this study.

PET/CT data were acquired with a GE Discovery STE (GE Healthcare, Waukesha, WI) using methods previously described. DW- and DCE-MRI were performed using a Philips 3T Achieva MR scanner (Philips Healthcare, Best, The Netherlands) using previously described methods. Following the DCE-MRI acquisition, a 3D T1-weighted high-resolution isotropic volume examination (THRIVE) scan was acquired. To improve the registration between the PET and MR data sets, a specially designed device that is an exact geometric replica of the breast support in the double-breast radiofrequency coil (In vivo Inc., Gainesville, FL) was constructed for PET/CT imaging.

2.B. Data analysis

While details of the DCE-MRI, DW-MRI, and FDG-PET analysis are presented in Refs. 15 and 16, the salient features are as follows. DCE-MRI data were analyzed using the extended Tofts-Kety relationship to yield estimates of the volume transfer constant (Ktrans), the extravascular extracellular volume fraction (ve), and the blood plasma fraction (vp), maps. The DW-MRI data were used to calculate the apparent diffusion coefficient (ADC) maps, and the FDG-PET data were used to calculate parametric maps of the FDG standard uptake value (SUV).

2.C. Image registration at each time point

Due to its relatively high spatial resolution and tissue contrast, we used the THRIVE images to align the PET and MR data sets acquired at each individual time point. This was achieved by using a rigid body registration (RBR) algorithm which searches for the optimal rotation and translational parameters by maximizing the normalized mutual information (NMI). The obtained transformations between the DCE-MRI and the THRIVE data, T RBR,DCE, at each time point (t1, t2, and t3), were directly applied to the Ktrans, ve, vp, and ADC maps at each time point to put all MRI parametric maps into a common image space. As the PET/CT

<table>
<thead>
<tr>
<th>Patients</th>
<th>Age (yr)</th>
<th>Treatment regimens</th>
<th>Receptor status</th>
<th>Tumor grade</th>
<th>Pathologic response</th>
<th>Excised tumor size (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>36</td>
<td>AC → taxol</td>
<td>ER + PR + HER2</td>
<td>2</td>
<td>Residual disease</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>48</td>
<td>Taxotere + carboplatin + herceptin</td>
<td>ER − PR + HER2</td>
<td>3</td>
<td>pCR</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>58</td>
<td>Cisplatin + taxol ± everolimus</td>
<td>ER − PR + HER2</td>
<td>2</td>
<td>Residual disease</td>
<td>1.7</td>
</tr>
<tr>
<td>4</td>
<td>33</td>
<td>AC → taxol</td>
<td>ER + PR + HER2</td>
<td>3</td>
<td>Residual disease</td>
<td>1.2</td>
</tr>
<tr>
<td>5</td>
<td>39</td>
<td>AC → taxol</td>
<td>ER − PR + HER2</td>
<td>3</td>
<td>pCR</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>43</td>
<td>Cisplatin + taxol ± everolimus</td>
<td>ER − PR + HER2</td>
<td>3</td>
<td>Residual disease</td>
<td>0.7</td>
</tr>
<tr>
<td>7</td>
<td>57</td>
<td>AC → taxol</td>
<td>ER − PR + HER2</td>
<td>3</td>
<td>Residual disease</td>
<td>N/A</td>
</tr>
<tr>
<td>8</td>
<td>55</td>
<td>AC → taxol</td>
<td>ER − PR + HER2</td>
<td>2</td>
<td>Residual disease</td>
<td>3.5</td>
</tr>
<tr>
<td>9</td>
<td>46</td>
<td>Taxotere + carboplatin + herceptin</td>
<td>ER + PR + HER2</td>
<td>3</td>
<td>Residual disease</td>
<td>0.3</td>
</tr>
<tr>
<td>10</td>
<td>39</td>
<td>AC → taxol</td>
<td>ER + PR + HER2</td>
<td>1</td>
<td>Residual disease</td>
<td>2.5</td>
</tr>
</tbody>
</table>

Medical Physics, Vol. 41, No. 5, May 2014
and THRIVE images were acquired on different scanners, the placement of the breast within each scanner resulted in different deformations. Thus, the CT data were first rigidly registered to the THRIVE images at each time point with the RBR, then a nonrigid body registration (NRBR) algorithm was applied. The RBR and the NRBR algorithms yielded the transformations $T_{RBR,CT}$ and $T_{NRBR,CT}$, respectively, which were applied to the SUV maps from the FDG-PET data.

2.D. Image registration across time

For a given patient, the two sets (or three sets for five of the patients) of THRIVE images were serially registered using the RBR and a modified version of the NRBR algorithm described above. In the modified algorithm, a constraint term was added to the NMI based cost function:

$$f_{\text{const}} = -NMI + \alpha \sum \left| \log(J_F(x)) \right|,$$

where $J_F(x)$ is the Jacobian determinant on the current voxel $x$, $M$ is the total number of voxels in the area, and $\alpha$ is the weight of the constraint term. For the patient data sets in this study we empirically chose $\alpha = 0.4$. This algorithm was designed to preserve the tumor volume while maximally registering surrounding tissues.

For all the patients, the THRIVE image at $t_1$ was registered to the THRIVE image at $t_2$ using both the RBR and the constrained NRBR algorithm to yield the transformations $T_{RBR,T1,T2}$ and $T_{NRBR,T1,T2}$, respectively. Similarly, the THRIVE image at $t_2$ was registered to the THRIVE image at $t_3$ to yield the transformation $T_{RBR,T2,T3}$ and $T_{NRBR,T2,T3}$, respectively, for the five patients whose data at $t_3$ were also available. To register the THRIVE images at $t_1$ to $t_3$, $T_{RBR,T1,T2}$, $T_{RBR,T1,T3}$, $T_{NRBR,T1,T2}$, $T_{NRBR,T1,T3}$, and $T_{NRBR,T2,T3}$ were then applied to the THRIVE images at $t_1$. Since the ADC, $K_{trans}$, $v_p$, and SUV maps at each time point were registered to their corresponding THRIVE image, these parametric maps could then be directly placed into a common image space by applying the transformation obtained from registering the THRIVE images longitudinally.

2.E. Registration validation

The tumor voxels in the SUV maps were determined by using a threshold of 40% of the maximum SUV uptake in the tumor ROIs (chosen based on previous reports), while the tumor in the DCE-MRI data was defined as the voxels within the outlined ROI that showed a postcontrast signal intensity that was $\geq 80\%$ (Refs. 27 and 28) of the precontrast signal intensity.

We qualitatively and quantitatively tested the validity of our approach. Qualitative assessment was performed by visual inspection of the alignment of the breast contours and tumors after registration. Quantitative assessment was done by calculating: (1) a modified Dice similarity index, (2) the bending energy, and (3) the change in the tumor volume after registration across time. Since the SUVs estimated from the FDG-PET data and the DCE-MRI data report on different aspects of tumor biology and moreover, different strategies were used to segment the tumor due to the large differences in the spatial resolutions in the PET and MR images, the original Dice similarity is not appropriate to compare two items of the same size. When two tumor regions are not in the same size, the largest possible Dice occurs when one segmented tumor region contains the other completed. Hence, we modified the Dice similarity measure to compare the degree of overlap between two objects (i.e., the tumor volumes as determined by PET and MRI) of difference sizes. That is, the modified Dice similarity is calculated as the original Dice similarity of the tumors between the SUV and the DCE-MRI images divided by the largest possible Dice:

$$Dice_{\text{new}} = \frac{2 \times n(A \cap B)}{n(A) + n(B)} \times \frac{2 \times \min(n(A), n(B))}{\min(n(A), n(B))},$$

where $A$ and $B$ are two regions and $n(\cdot)$ is the number of voxels in a region.

The bending energy describes the smoothness of the deformation field. The smoother the deformation field, the lower the bending energy. The bending energy and change in the tumor volume were calculated after applying the unconstrained and the constrained NRBR algorithms and compared using a Wilcoxon rank sum test to determine: (1) if the results returned by the two algorithms were significantly different and (2) if the algorithms significantly affected the tumor volume. This second point is of particular interest, as it is imperative that longitudinal registration minimally impact tumor size so that any changes in tumor size are due to biological changes and not those artificially induced by the registration algorithm.

3. RESULTS

The rows of Fig. 1 display the ADC, postcontrast DCE-MRI, THRIVE-MRI, CT, and the SUV of the FDG-PET images before (Columns A, B, and C) and after (D, E, and F) registration for a nonresponder at three time points. As demonstrated in the figure, there was poor alignment of the breast contours and the tumors before registration, but after registration there was excellent alignment of the breast contours and the tumors between the modalities at all imaging time points. Figure 2 shows the results of the registration of the parameter maps $K_{trans}$, $v_p$, ADC, and SUV of the FDG-PET maps (rows 1–5, respectively) superimposed on the anatomical THRIVE image at three time points for a representative patient. Note the (visually) excellent alignment which allows for examination of the variations in intratumoral spatial distributions of the various parameters.

The median of the modified Dice was 0.92. The medians of the percentage change in the tumor volumes from $t_1$ to $t_2$ were $-9.25\%$ and $-1.13\%$ for the unconstrained and
algorithms ($p = 0.0003$), respectively. Similarly, the medians from $t_2$ to $t_3$ were 0.0422 and 0.0000, respectively ($p = 0.0079$). Finally, the medians from $t_1$ to $t_3$ were 0.0973 and 0.0003, respectively ($p = 0.0079$). The median bending energy for all the time points was 0.0092 and 0.0001 for the unconstrained and constrained algorithms, respectively ($p = 2.84 \times 10^{-7}$).

4. DISCUSSION AND CONCLUSIONS

A survey of the literature reveals that the overwhelming majority of studies on breast registration focus on the registration of dynamic scans to correct for bulk motion that occurs during a single imaging session. There are only a few of studies\(^{11,12,30,31}\) that have made use of longitudinal registration of breast MRI data. However, these studies either did not focus on the registration method itself or did not account for tumor changes observed between scans. Our study is the first report of a method that enables the longitudinal registration of multimodal breast images and the corresponding parametric maps (i.e., $K_{\text{trans}}$, $v_e$, $v_p$, ADC, SUV) to a common space, thereby allowing for a more comprehensive analysis of tumor behavior at the voxel level. The results of this study are potentially of general interest as more efforts are made to synthesize multiparametric studies for predicting the response to breast cancer in the neoadjuvant setting. We showed good alignment between the images both by visual assessment and by the calculation of the intersection of the tumor volumes, the bending energy, and the change of tumor volumes. The error in the intersection was mainly due to the differences in positioning of the breast between the PET and the MRI acquisitions at each individual time point (i.e., $t_1$, $t_2$, or $t_3$). Another source of error is that the contrast between the tumor and normal tissues of low-dose CT images is much lower than the MR images collected in this study; this brings difficulties during the alignment between CT and MR. Ideally, we would want to achieve a registration error that is within the spatial resolution of the PET image.

One of the limitations of this study is that the parameter $\alpha$ is selected empirically and cannot (currently) be determined automatically. Moreover, the registration algorithm may not correct the change in the whole breast volume over time due to, for example, hormonal fluctuations over the menstrual cycle. Future technical efforts will focus on automating the optimal selection of this parameter, as well as employing external fiducial markers to improve the registration performance. Future clinical efforts will include identifying the spatial-temporal relationships between the parametric maps at the voxel level and integrating the relationships between parameter maps to optimize the prediction of the response of breast tumors to NAT. The aligned parameter maps can also be used to initialize and constrain mathematical models of tumor growth and treatment response.\(^{32}\)

ACKNOWLEDGMENTS

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