Methods, cont’d…

ADC volumes are computed from the interleaved b0 and b800 MRI DWI acquisitions. For each pair of registered ADC images a 128x128 joint density histogram (JDH) is constructed by incrementing the count of the 2D histogram defined by the two ADC values of the registered tumor. For the JDH of the two pre-Tx exams, inevitable sample bias is removed and variance is generalized, i.e. increased, by adding the transpose of its JDH to itself. Then a percentile threshold of the null distribution is estimated; here we demonstrate the use of the 97.5th percentile (Figure 2, top left).

Next, the same process is repeated using one of the short interval, pre-Tx exams and the post-Tx scan. This distribution represents the treatment effect (Figure 2, top right) and clearly reveals that ADC values have changed in several ways: first the mean has moved upwards, and secondly there are many more counts above the 97.5th percentile line derived from the null distribution. For each patient the first two rows of the table (Figure 2, middle) show the incremental percent increase in counts above the null’s 97.5th and 95th percentile for the Treatment Effect distribution.

At this stage increases above both the 97.5th and 95th percentile correlate perfectly with clinically assessed partial response (cPR), while decreases correlate perfectly with clinically assessed stable disease (cSD) for the first cycle of chemotherapy for these first 5 patients; the chances of our obtaining these 5 outcomes randomly is 3%. The bottom row of images in Fig. 2 shows a single slice of each the anatomical reference breast exams overlaid with red-green-blue mask of the tumor. Here red indicates the presence of voxels whose ADC changes (postTx minus preTx) are greater than the 97.5th percentile of the null distribution (regions of effect, i.e. cell kill, as well as limited noise); green indicates voxels whose changes are within the 2.5th – 97.5th percentiles (regions of no significant change) and blue indicates changes that are below the 2.5th percentile of the null (regions of effect, i.e. continued tumor growth, as well as limited noise). Responders shown in the yellow columns along with the %increase of counts above that expected from the null hypothesis. Non-responders shown in the orange columns demonstrate the corresponding decrease.

Recall that 8-11 days post-initiation of the first (AC) cycle of chemotherapy is very early in assessing tumor change compared with any other technique. Presumably these effect changes increase roughly proportionally to the time interval between the pre-therapeutic and post-therapeutic scans for several weeks. It is very encouraging indeed that we may reliably see changes within 8-11 days. Moreover for these five patients the 97.5th percentile corresponds to ADC changes whose mean is not significantly different than ±0.5 10−3 mm²/s, the same threshold used for the successful differentiation of PD and SD in glioblastoma multiforme brain cancer patients assessed with the similar methodology in our previous publications [1-3].

Methods

This is a double-blinded study where the blinded estimates of response to therapy have been obtained from registered voxel-wise apparent diffusion coefficient (ADC) scans which were then used to predict individual patient response to the first cycle of neoadjuvant chemotherapy. Likewise in a blinded fashion the breast oncologist independently estimated the clinical response of the patient at the end of the first adriamycin/cyclophosphamide (AC) phase before initiation of the second phase of therapy involving Taxotere.

In this protocol patients with breast CA that have elected neoadjuvant chemotherapy prior to surgery receive 2 baseline exams, typically within a 15 minute interval where the patient is removed from the scanner and then repositioned for the second scan; these short interval exams are used to observe a sample of the null change distribution since no macroscopic changes have occurred to the tumor in this interval. The initiation of the first cycle of chemotherapy (AC) is typically within one day of the short interval exams. Another MRI exam is obtained 8-11 days post-initiation. Tumor volumes of interest (VOI) were drawn on the anatomical image volume and were warped from the anatomical volume onto one of the two pretherapy (pre-Tx) diffusion volumes denoted as the reference. Subsequent registrations either between the two pre-Tx exams or the two pre- and post-Tx scans are also warped to account for repositioning deformations to the breast as well as any small compartmental changes to the tumor.

Warping is accomplished using thin plate splines where the degrees of freedom (DOF) of the warp is related to the local mutual information density and volume of the tumor. The user only picks the loci of 3 control points in the second tumor volume that approximates their loci in the reference tumor volume. The multiscale registration first implements rigid body registration, then low DOF warping, and finally full DOF warping [4].

References


Funded by NIH grants 1P01CA85878 & 1P01CA87634

![Figure 1: Registration of two interval breast exams, one shown in aqua hue and the other in grayscale.](image1)

![Figure 2: Five patients are evaluated for response. The yellow columns show responders (%counts above null threshold increased); non-responders shown in orange columns decreased.](image2)

![Figure 3: Adjacent slices from a registration of only the lesion in interval breast exams, one shown in yellow and the other in gray.](image3)