INTRODUCTION Non-Hodgkin’s lymphoma is the fifth most common cancer in the United States. During the last 50 years, its incidence has increased two-fold. The primary method of diagnostic imaging of this disease is by PET-CT, which detects the influx and retention of $^{18}$F-2-deoxyglucose by cancer cells, which is diminished after successful therapy. As PET requires administration of a radioactive tracer and fasting before imaging, we explored the use of more convenient and potentially less hazardous MRS and MRI methods for early detection of therapeutic response. In preclinical lymphoma studies in animal models, we demonstrated that $^1$H MRS and MRI provide sensitive markers for chemotherapy, immunotherapy and radiation therapy. Lactate and the apparent diffusion coefficient (ADC) are sensitive indices of response to chemotherapy and radiation therapy, while total choline is an indicator of response to immunotherapy and radiation therapy [1-3]. We have recently developed a Hadamard slice-encoded selective multiple quantum coherence sequence for lactate detection and implemented it on a clinical scanner [4]. The lactate imaging method and the single voxel PRESS and multi-slice diffusion-weighted imaging (DWI) sequences were utilized in this preliminary study of non-Hodgkin’s lymphoma patients.

METHODS The multi-slice lactate imaging sequence was previously demonstrated to produce high quality localized images of lactate in an NHL patient [4]. The sequence starts with a Hadamard encoded adiabatic slice inversion pulse for multi-slice selection. A selective multiple quantum coherence (Sel-MQC) pulse train follows thereafter; 90° (CH)$^3$ - 1/2J- 90° (CH) - 180° (CH$_3$) - 90° (CH$_3$)-1/2J-ACQ. Coherence selection gradients were inserted in the Sel-MQC pulse train between the second and third RF pulses and after the fourth RF pulse to select the double quantum to zero quantum transition of J-coupled lactate (CH$_3$) spins (1.3 ppm) while spoiling the signal of uncoupled lipid resonances resonating at the same frequency. 2D-CSI encoding gradients were placed after the last RF pulse. A Hadamard-slice encoded water spin echo CSI sequence was run for calibration. Total choline was measured using a single voxel PRESS sequence with outer volume suppression; TR=1500 ms, TE=135 ms. Unsuppressed water signals in the same voxel were used for calibration. The ADC was measured using a diffusion-weighted echo planar imaging (DW-EPI) sequence. B-values of 0, 250 s/mm$^2$, 500 s/mm$^2$, and 750 s/mm$^2$ were used. All measurements were performed on a 3T Siemens Trio system with the VB17 IDEA software. The Had-Sel-MQC-CSI raw data were retrieved using the TWIX command in the scanner and processed with an in-house IDL program. The PRESS spectral data were exported in .rda format and processed using NUTs software (Acorn NMR). ADC maps were generated using a Siemens-provided processing routine. MRS/MRI measurements were performed before treatment and within one week after initiation of treatment.

RESULTS Fig 1 shows data from a lymphoma patient with a lesion in the axillary node. Fig 1A, 1C and 1E are the T2-weighted image, ADC map and PRESS spectrum of the patient before treatment. Fig 1B, 1D and 1F are corresponding images and spectrum 4 days after initiation of therapy. An increase of ADC and decrease of tCho were observed. Quantitative changes appear in the Penn-3 row of Table 1. Table 1 summarizes the results of three non-Hodgkin’s lymphoma patients who participated in this preliminary longitudinal MRS/MRI study. The patient Penn-1 exhibited complete remission as observed at 6 month and continues at 2 years from treatment. Patient Penn-3 is in remission as of 5 month after treatment. The status of the patient Penn-2 is still being evaluated. The trend shows a decrease of Lac and tCho and increase of ADC after treatment. In one patient tCho increased rather than decreased. In view of our xenograft study [2], this may indicate that the patient is refractory to rituximab treatment, but further studies are required to confirm this conclusion.

DISCUSSION These initial studies of three NHL patients clearly demonstrate that MRS/MRI methods can detect early treatment responses within about 48 hr in non-Hodgkin’s lymphoma patients.

REFERENCES

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