Award Number: W81XWH-10-1-0743

TITLE: A Novel Multivoxel-Based Quantitation of Metabolites and Lipids Noninvasively Combined with Diffusion-Weighted Imaging in Breast Cancer

PRINCIPAL INVESTIGATOR: Michael Albert Thomas Ph.D.

CONTRACTING ORGANIZATION: University of California, Los Angeles
Los Angeles, CA 90024-1406

REPORT DATE: September 2012

TYPE OF REPORT: Annual

PREPARED FOR: U.S. Army Medical Research and Materiel Command
Fort Detrick, Maryland 21702-5012

DISTRIBUTION STATEMENT: Approved for Public Release; Distribution Unlimited

The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision unless so designated by other documentation.
**Purpose:**

- i) To extend the single-voxel based 2D MRS version of L-COSY to multi-voxel based analogue on a 3T MRI/MRS scanner using the echo-planar imaging (EPI) based spatial encoding for determining metabolic distributions over many voxels.

- ii) To implement a Matlab-based post-processing algorithm in order to process the 2D COSY data recorded in breast cancer.

- iii) To record DWI and to calculate ADC maps in breast cancer patients and healthy controls.

- iv) To correlate the changes in metabolite and lipid levels with ADC changes in breast cancer patients and healthy women.

**Scope:**

Improving the specificity of malignant and benign tumors will be a major outcome. Improved imaging techniques will enable unambiguous measurement of metabolites and the lipids in situ, which could potentially complement the existing diagnostic modalities commonly used in breast carcinoma.

**Major Findings:**

The MR protocol including 4D EP-COSI and DWI-MRI was successfully evaluated in six healthy women, 3 benign and 2 malignant breast cancer patients on a 3T MRI scanner. Report of the Progress: Multi-slice DWI-MRI and 4D EP-COSI was tested in 2 malignant and 3 benign breast cancer patients and 6 healthy women. The ADC maps using DWI and lipid/water images using EP-COSI have been derived. DWI images showed reduced ADC values in affected mass and the EP-COSI data showing lipid and metabolite changes in breast cancer.
# Table of Contents

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Introduction</td>
<td>4</td>
</tr>
<tr>
<td>Body</td>
<td>4</td>
</tr>
<tr>
<td>Key Research Accomplishments</td>
<td>10</td>
</tr>
<tr>
<td>Reportable Outcomes</td>
<td>10</td>
</tr>
<tr>
<td>Conclusion</td>
<td>11</td>
</tr>
<tr>
<td>References</td>
<td>11</td>
</tr>
<tr>
<td>Appendices</td>
<td>13</td>
</tr>
</tbody>
</table>
Introduction:

A leading cause of cancer deaths among women worldwide is due to breast cancer and new therapies and optimal treatments are continuously being conceived and explored to better control or even cure this disease. (1). Diagnosis and therapeutic management of the breast tumor remain significant medical challenges, hence early detection, diagnosis, and timely treatments are essential to successful health care (2). Magnetic resonance imaging and spectroscopy (MRI/MRS) have gained significant importance during the last fifteen years for the diagnosis and monitoring of breast cancer therapy. The sensitivity of MRI/MRS for anatomical delineation is very high and the consensus is that MRI is more sensitive in detection than x-ray mammography. Advantages of MRS include delivery of biochemical information about tumor metabolism, which can potentially assist in the staging of cancers and monitoring responses to treatment. High sensitivity is a major advantage of contrast enhanced MRI, but its diagnostic relevance in the future will largely depend on improvements in specificity. Current approaches in the application of MRI to breast tumors aim to improve specificity and sensitivity (4-17). Increased specificity is necessary to reduce the number of biopsies performed to confirm false positive findings. Diffusion-weighted imaging (DWI) is another MR based technique that probes the microstructure of tissues and is sensitive to the degree to which motion of water molecules is restricted in relation to how packed together cells are (17, 18). It has been reported that high resolution DWI may add valuable functional information to conventional MR protocols with short measurement times for the diagnosis of breast cancer and improve the specificity of MR imaging (19-21). However, new technological developments are necessary to assess their role in breast diagnosis. A method capable of identifying biochemical characteristics non-invasively in the tumor lesions that can be used in conjunction with MRI is proton (\(^1\)H) MR Spectroscopy (MRS). Researchers have shown that \(^1\)H MRS can be used to characterize breast cancers with improved diagnostic accuracy (22-26). Unfortunately, multi-voxel based novel MR spectroscopic imaging (MRSI) techniques using the speed advantage offered by echo-planar imaging (EPI) and improved spectral resolution offered by two-dimensional (2D) MR spectroscopy (MRS) have not been fully explored in breast cancer studies so far. Hence, a major task of this project is to combine echo-planar correlated spectroscopic imaging (EP-COSI) with DWI approach for improving the overall specificity of breast cancer detection.

Body:

i) Proposed Task 1: To further optimize the multi-voxel based extension of the correlated spectroscopy (COSY) sequence, in which two spectral encodings will be combined with two spatial encodings. This four-dimensional (4D) data acquisition scheme will be accomplished utilizing the echo-planar imaging (EPI) approach that is commonly used for spatial encoding in MRI including DWI. (Months 1-6).

This task was completed already (September 2010-May 2011) as reported last year.

ii) Proposed Task 2: To evaluate the EP-COSI data using a breast phantom containing two concentric spheres, the inner one containing several metabolites which have been reported in breast tissues surrounded by the outer phantom containing corn oil to mimic
fatty tissues known to be in breast tissues, and to optimize the echo speed-factor and other acquisition parameters using the phantom (Months 1-6).

This task was completed already (September 2010-August 2011) as reported again last year.

**iii) Proposed Task 3:** To develop, evaluate and optimize the prior-knowledge basis set spectra using the GAMMA-simulation and breast phantom solutions as prior knowledge for the multi-voxel based COSY spectra recorded using the 3T MRI scanner (Months 3-9).

Using the GAMMA library (27), a prior-knowledge basis-set containing metabolites and lipids was prepared (April 2010-August 2011) as reported last year.

**iv) Proposed Task 4:** To record the EP-COSI spectra in the fatty, glandular and ductal areas of healthy breasts. Twenty healthy female volunteers (25-70 years old) with no previous history of breast cancer will be investigated. (Months 6-24).

Five healthy women (age=36-59 years) were investigated using the DWI and EP-COSI MRI protocol as reported earlier (March 2011-July 2011). Six more healthy women (age=26-46 years) without any prehistory of breast cancer have been studied during the 2nd year. The ADC image derived from the DWI data recorded in a 34 yo healthy woman is shown in Fig.1. Feasibility of recording 2D COSY spectra in multiple regions using the EP-COSI sequence is demonstrated in Fig.2. Multi-voxel metabolite spectra of the fat peak at 5.3ppm is shown in Fig.2B (28). 2D MR spectra were recorded in multiple regions including the fatty and glandular regions of the 34 yo healthy subject as shown in Fig.2C and D. C and D were extracted COSY spectra from the fatty and glandular regions with a volume of 1x1x2cm³. These findings were reproduced in other 10 ten healthy women as shown in our earlier manuscript using a single voxel (SV) localized L-COSY sequence (29).

**Figure 1.** A) An axial ADC slice image recorded in a 34 yo healthy subject using the DWI sequence.
**Figure 2.** A) An axial MRI slice image recorded in the same healthy subject as above showing the EP-COSI localization; the small white box shows the regions of interest (ROI) localized the EP-COSI sequence and the large white box with 16x16 grids shows the spatial localization encoded by the EP-COSI sequence. B) Multi-voxel display of the EP-COSI data showing the olefinic diagonal peaks. C) and D) are 2ml 2D COSY spectra from the fatty and glandular regions extracted from the EP-COSI data.

**v) Proposed Task 5:** To record the multi-voxel-based 2D spectra in patients with benign and malignant breast cancer. The breast metabolite and lipid concentrations calculated from the multi-voxel data using the ProFit algorithm will be compared with LC-Model processed 1D spectral based MRSI data. Twenty patients with biopsy-proven breast cancer (ductal carcinoma and invasive lobular cancer), twenty patients with benign breast tumor (fibroadenoma, proliferative fibrocystic change and papillomas) will be investigated (Months 6-24).

Three women with benign (age=28-44 years) and 2 with malignant breast cancer (age=61 years) have been investigated using the 4D EP-COSI sequence on the 3T MRI scanner using a dedicated breast MRI coil. Shown in Fig. 3 are the following: A) An axial MRI slice image recorded in a 44yo benign cancer patient showing the EP-COSI localization. B) Multi-voxel EP-COSI spectra and C) a 2ml COSY spectrum recorded from the affected mass.
Figure 3. (A) An axial MRI slice image recorded in a 44 yo subject with benign breast cancer showing the EP-COSI localization; the small white box shows the regions of interest (ROI) localized the EP-COSI sequence and the large white box with 16x16 grids shows the spatial localization encoded by the EP-COSI sequence. (B) multi-voxel 2D COSY spectra covering fatty and glandular breast regions. (C) A 2ml benign lesion-indicating spectrum showing slightly increased water and slightly reduced lipids.
Figure 4. (A) An axial MRI slice image recorded in a 61 yo subject with malignant breast cancer (a hypo-intense mass) showing the EP-COSI localization; the small white box shows the regions of interest (ROI) localized the EP-COSI sequence and the large white box with 16x16 grids shows the spatial localization encoded by the EP-COSI sequence. (B) multi-voxel 2D COSY spectra covering fatty and glandular breast regions. (C) and (D) show 1ml spectra showing increased water and slightly reduced lipids.

Fig. 4 shows the following: A) An axial MRI slice image recorded in a 61yo malignant breast cancer patient showing the EP-COSI localization. B) Multi-voxel EP-COSI spectra; C) and D) show 2ml COSY spectra recorded from the affected mass and a
healthy region. These results are also in agreement with our earlier data on the SV-based L-COSY sequence (29-30).

**vi) Proposed Task 6:** To record multi-slice DWI in twenty patients with biopsy-proven breast cancer, twenty patients with benign breast tumor and twenty healthy women and to calculate the ADC maps. *(Months 6-24).*

**Figure 5.** An axial ADC slice image recorded in the 44 yo subject with a benign mass using the DWI sequence.

**Figure 6.** An axial ADC slice image recorded in the 61 yo subject with a malignant mass in the left breast using the DWI sequence.
The diffusion weighted imaging (DWI) has been recorded in three women with benign and 2 with malignant breast cancer 3T MRI scanner using a dedicated breast MRI coil.

vii) Proposed Task 7: To correlate the EP-COSI findings with that of DWI in differentiating benign from malignant breast cancers, and to calculate specificity, sensitivity and accuracy of the MRSI and DWI data in differentiating benign from malignant tumors. (Months 6-24).

This task will be completed during the next year.

Key Research Accomplishments

- The 4D EP-COSI has been successfully evaluated in 11 healthy women, 3 benign and 2 malignant breast cancer patients so far using the UCLA Radiology Siemens 3T MRI scanner equipped with a dedicated breast phased-array assembly. As summarized in the last year report, this sequence is available now at UCLA only and is not supplied by any of MRI manufacturers.
- The DWI-MRI protocol was successfully evaluated in a total of 11 healthy women, 3 benign and 2 malignant breast cancer patients so far.
- We have demonstrated that the EP-COSI spectroscopic imaging sequence can be combined with a clinical breast DWI protocol with the total duration of less than an hour. The protocol is safe enough to be included in any MRI protocol to be evaluated in breast cancer for improving the overall specificity.
- We are currently testing retrospective Maximum Entropy and Compressed Sensing of the 4D EP-COSI data so that the acquisition can be accelerated in the near future to 5-10 minutes which will lead to less inconvenience to patients suffering from breast cancer.

Reportable Outcomes:


B. Presentations: There were three conference presentations containing four abstracts.
C. Books: None on Breast Cancer Research based.

Conclusions: The scanning protocol including EP-COSI and DWI-MRI has been evaluated in 11 healthy women, three benign and 2 malignant breast cancer so far. We will continue to recruit 10 malignant, 10 benign and 5 healthy women during the next year. Our inability in recruiting more breast cancer patients has been discussed with the UCLA mammo co-investigators, namely Dr. Naneltte DeBruhl and Dr. Lawrence Bassett. They have promised us to improve the patient recruitment in the 3rd year.

References

Appendix:

A) A copy of our Poster presented at the at the International Society of Biomedical Engineering (ISBI) Conference 2012, Barcelona, Spain
MAXIMUM ENTROPY RECONSTRUCTION OF ECHO-PLANAR CORRELATED SPECTROSCOPIC IMAGING DATA OF HUMAN BREAST IN VIVO

Brian L. Burns and M. Albert Thomas
Departments of Biomedical Engineering and Radiological Sciences, David Geffen School of Medicine, University of California, Los Angeles, CA 90095

Introduction

• 4D Echo-Planar based Correlated Spectroscopic Imaging (EPCOSI) allows for the simultaneous acquisition of two spatial (k_y, k_x) and two spectral (t_2, t_1) dimensions in vivo in a single Magnetic Resonance (MR) experiment [1].

• The EPI readout accelerates the direct acquisition of the k_x and t_2 dimensions but the k_y and t_1 dimensions are incrementally acquired as indirect dimensions.

• Non-uniform under-sampling (NUS) the k_y and t_1 dimensions to 25% and reconstructing the missing data with Maximum Entropy (MaxEnt) preserves the spectral and spatial resolution of the acquired data and reduces the scan time from 20 to 5 minutes in healthy human breast in vivo.

Materials and Methods

• An EPCOSI scan of healthy breast with the following parameters: 1x1x2cm^3 voxel size, TR/TE=1.5s/30ms, 50 t_1 increments, 16x16cm^2 FOV was retrospectively NUS to 25% of the k_y-t_1 plane using the mask in Fig. 2. Figure 1 (Top) Selected 2D COSY spectra from a 4D EPCOSI scan of healthy, fatty breast highlighted in red in the MRI localizer image for A1) fully sampled B1) 25% NUS C1) 25% NUS with MaxEnt reconstruction. (Bottom) Spatial distribution of olefinic cross-peaks of unsaturated fatty acids highlighted in light blue for A2) fully sampled B2) 25% NUS C2) 25% NUS with MaxEnt reconstruction.

• Missing data in the k_y-t_1 plane was reconstructed using MaxEnt [2,3]:

\[
\maximize S_{1/2}(f) \text{ s.t. } C(f) \leq \sigma
\]

where \( f \) is the estimated fully-sampled spatially distributed spectra, \( C(f) \) is the time-domain fidelity constraint, \( \sigma \) is the noise standard deviation, and \( S_{1/2}(f) \) is the spin-\( 1/2 \) entropy of the estimated spectra [4].

• The Cambridge MaxEnt algorithm uses a variant of the conjugate gradient method to solve (1) as shown in Fig. 3.

Results and Discussion

• A1 is a voxel from a fully sampled 4D EPCOSI scan of healthy, fatty breast in vivo. B1 is the same voxel after 25% NUS of the k_y-t_1 plane and has significant artifacts from the large diagonal peak at (2.2ppm, 2.0ppm) obscuring the much smaller cross-peak highlighted in blue. C1 shows the MaxEnt reconstruction of B1 and has equivalent spectral peaks to the fully sampled data in A1 with the artifacts removed.

• A2, B2, and C2 show the spatial distribution of the olefinic cross peaks highlighted in blue for the fully sampled, 25% NUS and 25% NUS with MaxEnt reconstruction from a 4D EPCOSI scan of healthy, fatty breast. B2 shows significant spatial incoherent artifacts from the NUS that are removed by MaxEnt reconstruction in C2.

• The normalized Root Mean Square Error (nRMSE) of B1/B2 is .0047 and .0014 for C1/C2.

Conclusion

• We have shown it is possible to simultaneously NUS a spectral and a spatial dimension of an in vivo EPCOSI scan to 25% and successfully reconstruct the data. This acceleration translates into a clinically viable 5 minute EPCOSI scan time.

Acknowledgements

• DoD Idea Expansion grant for Breast Cancer Research #W81XWH-10-1-0743 and NIH training grant #ST1 LM07356
Maximum Entropy Reconstruction of Correlated Spectroscopy of Human Breast In Vivo

Brian L. Burns1,2, Jon K Furuyama1, Neil Wilson1, Nicki DeRuhl1, Alex Bu12, and M. Albert Thomas3
1Biomedical Engineering, UCLA, Los Angeles, CA, United States, 2Medical Imaging Informatics (MII), UCLA, Los Angeles, CA, United States, 3Department of Radiological Sciences, UCLA

Introduction – The altered metabolism of breast cancer tissues gives rise to changes in metabolite concentrations that are detectable using two-dimensional (2D) Magnetic Resonance Spectroscopy (MRS) [1]. 2D Localized-Correlated Spectroscopy (L-COSY) MRS has been shown to increase the specificity and sensitivity of tumor grade classification when used with traditional techniques, such as Dynamic Contrast Enhanced (DCE)-MRI [1]. However, because of the number of t₁ increments used to construct the indirect spectral dimension (F₂), the L-COSY sequence requires on the order of 25 minutes for 100 t₁ increments and 8 averages [2]. To reduce scan times to a clinically acceptable 12 minutes, only 45 t₁ increments are typically used, but this reduces spectral resolution along F₁. Maximum entropy (MaxEnt) image reconstruction techniques have been used in Nuclear Magnetic Resonance (NMR) to non-uniformly under-sample (NUS) indirect spectral dimensions and then reconstruct the most statistically likely fully-sampled multi-dimensional spectrum [3,4]. This technique can be used to accelerate the collection of 2D L-COSY data in vivo by selectively under-sampling along t₁. We show that under-sampling the indirect dimension by ~50% and reconstructing the spectrum using MaxEnt preserves the metabolite diagonal peaks: olefinic fat (UFD), methyl fat (FMETD), and choline (Cho) and the cross peaks: unsaturated fatty acid, right (UFR), unsaturated fatty acid, left (UFL), and triglycerol fat (TGFR) which have demonstrated clinical importance.

Methods – To qualitatively measure the performance of the MaxEnt reconstruction algorithm on NUS in vivo data, fully sampled 2D L-COSY scans of both healthy (n=10) and malignant (n=8) breast tissues were acquired on a 3T Siemens Trio scanner using the following parameters: 1cm³ voxel size, TR/TE=2s/30ms, 45 t₁ increments, 8 averages, and no water suppression. Both scans were retrospectively under-sampled to 22 lines along the t₁ dimension, apodized along the t₁ and t₂ dimensions, and then reconstructed using the MaxEnt algorithm to 45 lines. MaxEnt is a constrained convex optimization algorithm that uses a variant of the conjugate gradient method to iteratively solve the inverse problem [5]:

\[
\text{maximize } S_{1/2}(f) \text{ s.t. } ||F^{-1}Kf - D||_2 \leq \sigma
\]

where \(f\) is the estimated fully-sampled spectrum at each iteration, \(F^{-1}\) is the inverse Fourier transform, \(K\) is the NUS matrix, \(D\) is the time-domain measured data, \(\sigma\) is the noise standard deviation, and \(S_{1/2}(f)\) is the spin-½ entropy of the estimated spectrum [3]. The spectrum with the highest entropy is that which conforms to the uniform distribution and has a flat baseline. Therefore, by maximizing the spin-½ entropy in (1), MaxEnt enforces sparsity of the estimated spectrum in the frequency domain and reduces incoherent aliasing artifacts in the spectrum caused by NUS. The fidelity constraint ensures that peaks in the estimated spectrum must come from the sampled data to within a tolerance of the noise. The reconstruction is performed over both dimensions simultaneously as opposed to a series of 1D reconstructions as implemented elsewhere [4].

Results and Discussion – Figures 1A and 1B show the 2D L-COSY spectra of the fully sampled and ~50% MaxEnt reconstruction of the NUS healthy breast tissue, respectively. As can be seen, the major diagonal and cross peaks present in healthy fatty breast tissue are present in both spectra. The peak locations, line shapes, and amplitudes of the NUS data are roughly equivalent to the fully sampled data; however a slight loss in spectral resolution along F₁ is evident by the elongation of the UFR peak in the MaxEnt reconstruction. There is no corresponding loss in resolution along F₂. Figures 2A and 2B show 2D L-COSY spectra of the fully sampled and the ~50% reconstruction of the NUS malignant fatty breast tissue, respectively. The presence of water in fatty breast tissue is an indicator of breast cancer, and because of its high concentration, its broad line-width under-sampling along t₁. We show that under-sampling the indirect dimension by ~50% and reconstructing the spectrum using MaxEnt preserves the metabolite diagonal peaks: olefinic fat (UFD), methyl fat (FMETD), and choline (Cho) and the cross peaks: unsaturated fatty acid, right (UFR), unsaturated fatty acid, left (UFL), and triglycerol fat (TGFR) which have demonstrated clinical importance.

Conclusions – We have shown that it is possible to undersample the L-COSY sequence by up to ~50% and reconstruct in vivo spectra from as few as 22 t₁ lines that show similar spectral characteristics to spectra from fully sampled data. This acceleration translates into a sub-6 minute scan time if 8 averages are used and increases the potential use of MRS for breast cancer screening in the clinic. If increased spectral resolution is needed, the MaxEnt reconstruction is not limited to 45 t₁ points and can reconstruct many more NUS points than what was used in these experiments.


Acknowledgements – DoD Idea Expansion grant for Breast Cancer Research #W81XWH-10-1-0743 and NIH training grant #5T15 LM07356
Maximum Entropy Based Reconstruction of Echo-Planar Correlated Spectroscopic Imaging of Human Breast In Vivo

Introduction – 2D Localized Correlated Spectroscopy (L-COSY) has been shown to be a powerful tool in the detection of breast cancer but is limited to a single voxel [1]. The recently introduced Echo-Planar based Correlated Spectroscopic Imaging (EPCOSI) sequence allows for the simultaneous acquisition of two spatial (k_y, k_x) and two spectral (t_2, t_1) dimensions in a single experiment [2]. The 4D EPCOSI sequence interleaves the acquisition of the k_y and t_1 dimensions along the EPI read-out, while k_x and t_2 lines are incrementally acquired as indirect dimensions, requiring 20 minutes for a typical scan. To reduce scan times using standard Fast Fourier Transform (FFT) based reconstruction techniques would require the reduction of either the k_x spatial or t_2 spectral dimensions, and a corresponding reduction in resolution. Maximum entropy (MaxEnt) image reconstruction techniques have been used in Nuclear Magnetic Resonance (NMR) to non-uniformly under-sample (NUS) indirect spectral dimensions and in imaging to NUS spatial dimensions then reconstruct the fully-sampled multi-dimensional spectra or image [3,4,5]. This technique can be used to accelerate the collection of 4D EPCOSI data in vivo by selectively under-sampling along k_x and t_2. We show that under-sampling the indirect dimensions by ~25% and reconstructing the spatially localized EPCOSI spectra using MaxEnt preserves the spatial distribution of the metabolite diagonal peaks found in healthy breast tissue and reduces the scan time to 5 minutes.

Methods – An EPCOSI scan of healthy breast was performed on a 3T Siemens Trio scanner with the following parameters: 1x1x2cm^3 voxel size, TR/TE=1.5s/30ms, 50 t_1 increments, 16x16cm^2 FOV, 1 average with water suppression and 2 averages without water suppression. The scan was retrospectively under-sampled to ~25% of the k_y-t_1 plane, using the map in Fig. 1. The NUS spatial and spectral dimensions were simultaneously reconstructed using MaxEnt. The MaxEnt algorithm uses a variant of the conjugate gradient method to iteratively solve the constrained convex optimization problem [5]:

$$\text{maximize } S_{1/2}(f) \text{ s.t. } ||F^{-1}Kf - D||_2 <= \sigma$$

where $f$ is the estimated fully-sampled spatially distributed spectra at each iteration, $F^{-1}$ is the inverse Fourier transform, $K$ is the NUS matrix, $D$ is the time-domain measured data, $\sigma$ is the noise standard deviation, and $S_{1/2}(f)$ is the spin-1/2 entropy of the estimated spectra [3]. The spatially distributed spectra with the highest entropy are those which conform to the uniform distribution and have a flat baseline. Therefore, by maximizing the spin-1/2 entropy in (1), MaxEnt enforces sparsity of the estimated spectra in the frequency and k-space domains and reduces incoherent aliasing artifacts in the spectra caused by NUS. The fidelity constraint ensures that peaks in the estimated spectra and their spatial distribution must come from the sampled data to within a tolerance of the noise. The reconstruction is performed over all dimensions simultaneously as opposed to a series of 1D reconstructions as implemented elsewhere [5].

Results and Discussion – Figures 2A and 2B show the fully sampled and 25% MaxEnt reconstruction of the selected 2D COSY spectra from a voxel of healthy fatty breast tissue. As can be seen, the spectral characteristics of the NUS data with MaxEnt reconstruction are equivalent to the fully sampled data; the same diagonal and cross peaks are present in both spectra without a significant increase in noise. Figures 3A and 3B show the spatial distribution of the olefinic cross peak selected in Figure 2 for the fully sampled and 25% MaxEnt reconstructed 4D EPCOSI of healthy fatty breast tissue. The spatial distributions between the fully sampled and MaxEnt reconstruction show very good agreement, with the exception of some spatial bleeding on the outer edges of the distribution.

Conclusions – We have shown that it is possible to under-sample both the spectral and spatial dimensions of an in vivo EPCOSI sequence by up to 25% and reconstruct the spectra with similar spatial distributions and spectral characteristics to the fully sampled data. This acceleration translates into a 5 minute EPCOSI scan time and increases the potential use of 4D MRS in the clinic. Compressed Sensing (CS)-based reconstruction can be use alternatively for processing the NUS EPCOSI data. Further research regarding the merits and demerits of the two reconstruction methods is in progress.


Acknowledgements – DoD Idea Expansion grant for Breast Cancer Research #W81XWH-10-1-0743 and NIH training grant #5T15 LM07356
# Maximum Entropy Reconstruction of Non-Uniformly Under-Sampled Multidimensional Spectroscopic Imaging in vivo

<table>
<thead>
<tr>
<th>Journal:</th>
<th>Magnetic Resonance in Medicine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manuscript ID:</td>
<td>MRM-12-13731</td>
</tr>
<tr>
<td>Wiley - Manuscript type:</td>
<td>ISMRM Young Investigator Award</td>
</tr>
<tr>
<td>Date Submitted by the Author:</td>
<td>11-Oct-2012</td>
</tr>
<tr>
<td>Complete List of Authors:</td>
<td>Burns, Brian; UCLA School of Medicine, Biomedical Engineering Wilson, Neil; David Geffen School of Medicine, Radiology Furuyama, Jonathan; David Geffen School of Medicine, Radiology Thomas, M. Albert; David Geffen School of Medicine, Radiology</td>
</tr>
<tr>
<td>Research Type:</td>
<td>Reconstruction &lt; Technique Development &lt; Technical Research, Spectroscopy/Spectroscopic Imaging &lt; Technique Development &lt; Technical Research</td>
</tr>
<tr>
<td>Research Focus:</td>
<td>No specific tissue or organ focus</td>
</tr>
</tbody>
</table>
Maximum Entropy Reconstruction of Non-Uniformly Under-Sampled Multidimensional Spectroscopic Imaging in vivo

Brian Burns¹,², Neil E. Wilson¹,³, Jon K. Furuyama¹,³, M. Albert Thomas¹,²,³

¹ Department of Radiological Sciences, David Geffen School of Medicine, University of California, Los Angeles, California, USA
² Department of Biomedical Engineering, University of California, Los Angeles, California, USA
³ Biomedical Physics IDP, University of California, Los Angeles, California, USA

Running Title: MaxEnt Reconstruction of Under-sampled MRSI Containing 2D Spectral Combined with 2D Spatial in vivo

Word Count: 5333

Address for Correspondence:
M. Albert Thomas Ph.D.
Radiological Sciences
David Geffen School of Medicine at UCLA
10833 Le Conte Avenue
Los Angeles, CA 90095-1721
Tel: (310) 206 4191
Fax: (310) 825 5837
Email: athomas@mednet.ucla.edu
Abstract

**Purpose:** To reduce scan times of 4D Magnetic Resonance Spectroscopic Imaging by applying non-uniform under-sampling (NUS) to the indirectly acquired dimensions ($k_y$ and $t_1$) and reconstructing those dimensions using Maximum Entropy (MaxEnt) iterative reconstruction.

**Methods:** Simulated and in vivo 4D Echo-Planar based Correlated Spectroscopic Imaging (EP-COSI) data were retrospectively under-sampled, and in vivo 4D Echo-Planar J-Resolved Spectroscopic Imaging (EP-JRESI) prostate data was prospectively undersampled in the $k_y$-$t_1$ plane. The under-sampled data sets were reconstructed using the Cambridge iterative algorithm to solve the MaxEnt reconstruction problem.

**Results:** Simulated data reconstructions demonstrated the MaxEnt reconstruction improves data fidelity considerably over the under-sampled data at SNRs from 2 to 20 and percent under-samplings of 20 to 80%. 25% NUS was applied retrospectively to EP-COSI and prospectively to EP-JRESI in vivo data. The NUS EP-COSI and EP-JRESI in vivo data were under-sampled to 25% and the incoherent artifacts caused by the under-sampling were successfully removed by the MaxEnt reconstruction.

**Conclusion:** MaxEnt reconstruction preserved the spectral-spatial resolution of the non-uniformly under-sampled data and reduced EP-JRESI scan times from 24 to 6 minutes in human prostate and can potentially reduce EP-COSI scan times from 20 to 5 minutes in human breast.

**Keywords:** EP-COSI, EP-JRESI, Maximum Entropy, Non-Uniform Under-Sampling, Spectroscopy, Spectroscopic Imaging
INTRODUCTION

The altered metabolism of cancer gives rise to changes in metabolite concentrations that can potentially be detected non-invasively using one-dimensional (1D) magnetic resonance spectroscopy (MRS) in vivo [1,2,3,4,5]. However, the overlap of spectral peaks in 1D MRS is a major concern for identifying individual metabolites. Two dimensional (2D) MRS has increased spectral dispersion over 1D MRS and can disentangle overlapping complex spectral peaks [6]. Single voxel 2D MRS has been shown to increase the specificity and sensitivity of tumor grade classification when used with traditional techniques, such as dynamic contrast enhanced (DCE)-MRI in the breast [7] and the Gleason grading system in the prostate [8]. However, requiring multiple $t_1$ increments per voxel to form the second spectral dimension limits its ability to provide multi-voxel coverage due to long scan times needed to combine two spectral and two spatial dimensions.

With the advent of Echo-Planar Spectroscopic Imaging (EPSI), magnetic resonance spectroscopic imaging (MRSI) scans with one spectral dimension could be completed within clinically acceptable times by interleaving the acquisition of a spatial and spectral dimension within the EPSI readout [9] [10] [11]. The recently introduced Echo-Planar based Correlated Spectroscopic Imaging (EP-COSI) [12] and Echo-Planar J-Resolved Spectroscopic Imaging (EP-JRESI) [13] sequences allow for the simultaneous acquisition of two spatial ($k_y, k_x$) and two spectral ($t_2, t_1$) dimensions in a single recording to form 4D MRSI. Both sequences interleave the acquisition of the $k_x$ and $t_2$ dimensions within the EPSI read-out, but $k_y$ and $t_1$ are incrementally acquired as indirect dimensions during each TR. The EP-COSI and EP-JRESI sequences have the benefits of increased spectral dispersion and multi-voxel support; however their scan times are directly proportional to the number of increments in the $k_y$ and $t_1$ dimensions and can take 20 to 40 minutes using typical parameters which is too long to be used for a routine clinical protocol [12] [13].

When using the Fast Fourier Transform (FFT), reducing 4D EP-COSI and EP-JRESI scan times requires the reduction of either the $k_y$ spatial or $t_1$ spectral dimensions through truncation or lower sampling rates, and a corresponding unwanted reduction in resolution.
or bandwidth with the potential for aliasing. However, non-uniform under-sampling of the spatial-spectral \( k_y-t_1 \) plane in combination with iterative non-linear reconstruction techniques can be used to accelerate the collection of 4D MRSI data in vivo while preserving the spatial and spectral resolution [14].

Earlier work has demonstrated the feasibility of under-sampling the mixed-domain \( k_y-t_1 \) plane of a 4D EP-JRESI data set and reconstructing the missing points with Compressed Sensing (CS) [14], a popular method of under-sampled iterative image reconstruction that promotes data sparsity in the reconstruction domain and data fidelity in the sample domain [15] [16]. The nature of spatial-spectral incoherent artifacts in the \( k_y-t_1 \) plane were explored, and it was shown that \( l_1 \)-norm-based CS reconstruction is a viable means of reducing the scan times of 4D EP-JRESI in vivo through under-sampling. In recent years, CS reconstruction has been successfully applied to under-sampled MRI [17] [18], 3D MRSI [19], dynamic MRI [20] [21], and a number of other promising applications not necessarily related to medical imaging [22] [23].

Maximum Entropy (MaxEnt) image reconstruction is an alternative non-linear iterative reconstruction technique to CS. It promotes the entropy of the data in the reconstruction domain while preserving data fidelity in the sample domain [24,25]. MaxEnt has been successfully applied to under-sampled images in astronomy and multi-dimensional spectra in NMR [25] [26] [27]. However, MaxEnt has not been applied to the mixed-domain \( k_y-t_1 \) plane of a 4D MRSI data set in vivo.

In information theory, entropy is a measure of the uncertainty of a variable state and the probability of its occurrence; the more uncertain its state, the higher its entropy. States that are all equally likely and whose probabilities conform to the uniform distribution have the maximum possible entropy [28]. Entropy in MRSI represents a measure of the phase coherence of an ensemble of spins within a given voxel volume [26]. Normalized for the concentration of spins within the volume, high-amplitude NMR signals represent states of full phase coherence and low entropy, while low amplitude NMR signals represent states of low phase coherence and high entropy. If the phase distribution
conforms to a state of low phase coherence, such as a uniform distribution, the entropy is maximized as it is in information theory. Therefore, by maximizing the entropy of the spatial, spectral-domain in MaxEnt, under-sampling artifacts are removed from the reconstruction because they represent states of high phase coherence and low entropy that are not present in the sampled data.

In this paper, we show that MaxEnt reconstruction is a viable technique to reduce scan times by successfully reconstructing under-sampled 4D MRSI data. We quantitatively characterize the MaxEnt reconstruction by showing results for retrospectively under-sampled simulated 4D EP-COSI data at varying levels of SNR and percent undersampling. Additionally, we show that under-sampling the k_y-t_1 plane to 25% and using MaxEnt to reconstruct an in vivo 4D EP-COSI data set retrospectively and an in vivo EJPRESI data set prospectively preserves the spatial-spectral resolution of the spectra and reduces scan times to 5 minutes in the breast and 6 minutes in the prostate using typical parameters.

METHODS

MR Simulations

The effects of SNR and percent under-sampling on the MaxEnt reconstruction were quantitatively assessed using a noise-free simulated 4D EP-COSI data set that contained either N-acetylaspartate (NAA), creatine (Cr), choline (Cho), lactate (Lac), or nothing in each voxel, as represented in the top of Figure 1 by a diagonal peak from each metabolite. Each metabolite was simulated using the GAMMA NMR libraries [29] from a Localized 2D Correlated Spectroscopy (L-COSY) sequence [30] with the following parameters: 100 t_1 increments, 1024 points in t_2, TR/TE= 1.5s/30ms, and spectral bandwidths of 1250Hz and 2000Hz along F_1 and F_2, respectively. Each 2D spectrum was line-broadened by 1 Hz and apodized by a sine-squared filter along t_1 and a skewed sine-squared filter with skew parameter 0.5 along t_2. They were then copied into an 8x8 spatial grid to simulate spatially distributed metabolites as follows: the upper left quadrant contained 2x2 voxels of NAA, the upper right quadrant contained 2x2 voxels of Cr, the lower left quadrant contained 2x2 voxels of Cho, and the lower right quadrant contained 2x2 voxels of Lac.
As a result of under-sampling the $k_y$-$t_1$ plane, the spatial-spectral artifacts caused the NAA and Cho voxels to alias into each other and the Cr and Lac voxels to alias into each other, as shown in the bottom of Figure 1. This changed the NAA to Cho and Lac to Cr ratios as the spatial and spectral separation between the metabolites broke down.

The simulated 4D EP-COSI data set was retrospectively reconstructed using MaxEnt at 20, 40, 60 and 80% under-sampling and SNRs of 2 to 20 in increments of 2. The SNR was varied by simulating different levels of thermal noise in the data set by adding Gaussian noise in quadrature to the noise-free 4D EP-COSI data set as in [31]. The desired SNR was achieved by ensuring the Root Mean Square (RMS) of the additive noise was equal to $1/SNR$ of the noise-free data set RMS, such that:

$$\text{noisy data} = \text{noise free} + \frac{\text{RMS(Noise Free)}}{\text{RMS(Noise)} \times \text{SNR}} \times \text{Noise}$$

[1]

Because the additive noise was random, each SNR was simulated and reconstructed 20 times per sample mask to account for random fluctuations in the reconstruction. The sampling masks were created using the 2D Poisson-gap method described below.

**MR Spectroscopic Imaging**

A 4D EP-COSI scan of a 34 year old healthy human breast was acquired on a Siemens 3T Trio scanner with the following parameters: 1x1x1 cm$^3$ voxel size, 50 $t_1$ increments, TR/TE/averages = 1.5s/30ms/1, a 16x16cm$^2$ FOV, and spectral bandwidths of 1250Hz and 1190Hz along $F_1$ and $F_2$, respectively. The scan was fully sampled and took 20 minutes to complete. Prior to being under-sampled and reconstructed, coil combination and eddy current correction were applied to the fully sampled data set. It was then apodized by a sine-squared filter along $t_1$ and a skewed sine-squared filter with skew parameter 0.5 along $t_2$. The fully sampled data set was then retrospectively under-sampled to 25% in the $k_y$-$t_1$ plane using the mask shown in Figure 4 and reconstructed by MaxEnt.
The prospective MaxEnt reconstruction was performed on a 4D EP-JRESI scan of a 71-year-old human prostate with cancerous tumors in the left and right base (Gleason score of 3+3, prostatic specific antigen of 8). The scan was acquired on a Siemens 3T Trio scanner using the single-channel endorectal coil with the same parameters as the EP-COSI breast scan except 64 t\textsubscript{1} increments and a spectral bandwidth of ±500Hz along F\textsubscript{1} was used. The k\textsubscript{y}-t\textsubscript{1} mask shown in Figure 5 was used during the scan to acquire 25% of the k\textsubscript{y}-t\textsubscript{1} plane during the acquisition. Eddy current correction was applied to the 4D prostate data after MaxEnt reconstruction.

Sample Mask Generation

The 2D Poisson-gap sample masks used were derived from 1D Poisson-gap sample masks [32]. Poisson-gap sample masks avoid large gaps between samples, which are detrimental to the reconstruction, while ensuring the samples are randomly distributed [33].

The effects of the sample mask on the SNR and line-shape of spectral reconstructions are well documented [33] [34] [35]. Sample masks that follow the time-domain NMR signal envelope and sample more points at higher SNR have lower RMS errors (RMSE) and non-linearity compared to sample masks that do not. Because the t\textsubscript{1} dimension in a filtered EP-COSI scan has a sine-bell signal envelope and the t\textsubscript{1} dimension in an EP-JRESI scan has an exponentially decaying signal envelope, the 2D Poisson-gap sample masks were modulated along t\textsubscript{1} with sine or exponential decay functions, respectively. The k\textsubscript{y} dimension for both scan types was modulated by an exponential decay function similarly to what has been used previously to maximize spatial SNR [36].

The rate parameter, $\lambda$, determines both the mean and variance of a Poisson distribution. The probability of generating a sample gap, $g$, from a Poisson distribution is characterized by:

$$p(g, \lambda) = \frac{\lambda^g \cdot e^{-\lambda}}{g!}$$  \hspace{1cm} [2]
For large $\lambda$, large values of $g$ are more likely, and for small $\lambda$, small values of $g$ are more likely. Therefore, the probability of $g$ can be modulated by varying the value of $\lambda$ according to a sine or exponential decay function, and the probability of small gaps can be increased where the SNR is highest in the MR signal envelope [32]. To generate $g$ as a function of $\lambda$, a Poisson process can be simulated using various techniques that do not depend on a priori knowledge of $g$ as above [37]. For these experiments, the poissrnd($\lambda$) function in Matlab was used to generate $g$ as a function of $\lambda$. It takes as input an array of $\lambda$ and returns an array of sample gaps with local mean and variance, $\lambda$.

An example of the 2D $\lambda$, gap, and mask arrays generated by Poisson-gap mask creation are shown in Figure 2 and illustrates their relationships; as the size of $\lambda$ and the gaps increase, the sample density decreases in that area of the mask. The magnitude Point Spread Function (PSF) of each mask is shown and demonstrates the viability of this approach; the single dominant central peak with small side-lobes, surrounded by low amplitude, incoherent artifacts is the desired profile of a random under-sampling mask PSF [38] [39].

**4D MaxEnt Reconstruction**

MaxEnt image reconstruction was used in these experiments to reconstruct the under-sampled data sets. MaxEnt reconstruction maximizes the entropy of the data in the spatial, spectral-domain while maintaining data fidelity with the sampled data in the k-space, time domain [25].

MaxEnt image reconstruction of 4D MRSI data is formulated as a constrained convex optimization problem [25] [27]:

$$\begin{align*}
\text{maximize } & S(m)^{1/2} \\
\text{subject to } & \|K F m - d\|_2^2 \leq C_0
\end{align*}$$

(3)
where \( m = (y, x, F_2, F_1) \) is the reconstructed spatial, spectral-domain data, \( F \) is the 4D Fourier operator, \( K \) is the under-sampling mask that determines which samples were acquired in the \( k_y-t_1 \) plane, \( d = (k_y, k_x, t_2, t_1) \) is the k-space, time-domain sampled data, \( C_0 \) is the standard deviation of the noise in \( d \), and \( S(m)^{1/2} \) is the entropy of the estimated spectrum.

The entropy used in these experiments was not the often used \(-\sum p \log(p)\) entropy introduced by Shannon [28] but the \( S_{1/2} \) entropy derived by Daniell and Hore specifically for NMR spectra originating from spin \( \frac{1}{2} \) nuclei, such as \( ^1H \) used in MRSI [26]:

\[
S(m)^{1/2} = -\sum \frac{|m|}{\text{def}} \log \left( \frac{|m|/\text{def} + \sqrt{4 + (|m|/\text{def})^2}}{2} \right) - \sqrt{4 + \left(\frac{|m|}{\text{def}}\right)^2}
\]  

(4)

where \( \text{def} \) is a scaling parameter related to the sensitivity of the scanner and is calculated for \( m \) of length \( N \) as \( \sqrt{C_0/N} \) [27]. The underlying physical processes that produce an MR spectrum are not based on particle, or discrete events so they cannot be modeled by simple Poisson processes as required for the derivation of Shannon entropy [40]. They are governed by the density matrix of the spin system under investigation. This equation can be applied to any MR spectrum originating from spin \( \frac{1}{2} \) nuclei and addresses previous concerns regarding the use of entropy in MRS and MRI reconstruction [41].

The MaxEnt reconstruction problem was solved by a Matlab implementation of the Cambridge algorithm [25]. It recasts MaxEnt image reconstruction into an unconstrained convex optimization problem and uses a variant of the conjugate gradient method to iteratively find the extrema in two phases; the first phase minimizes the fidelity constraint, and the second phase maximizes the entropy, while keeping the fidelity constraint minimized. The stopping criterion for the problem is reached when the gradients of the entropy, \( S(m) \), and fidelity constraint, \( C(m) \), are parallel: 

\[
\frac{\nabla S}{||\nabla S||_2} - \frac{\nabla C}{||\nabla C||_2} < 
\]
Specific details on the algorithm and modifications to accommodate multidimensional MR data can be found in [27].

RESULTS

MR Simulations

Quantitative results for the MaxEnt reconstruction of the simulated 4D EP-COSI data set at different percent under-samplings and SNR are shown in Figure 3. As described earlier, noise was added to the noiseless 4D data set in order to simulate SNRs of 2 through 20 in increments of 2 which were then undersampled to 80%, 60%, 40%, and 20% and reconstructed by MaxEnt. This was repeated 20 times per SNR and percent under-sampling in order to produce a mean and standard deviation for each metric.

The left pane of Figure 3 shows the average RMSE versus SNR for the under-sampled and MaxEnt reconstructed data sets at each percent under-sampling. The RMSE provides an estimate of reconstruction accuracy with respect to a reference data set that increases as the two data sets become more dissimilar. The RMSE was calculated in the spatial-spectral domain as:

\[
RMSE = \frac{1}{N} \sqrt{\sum (|\text{data}| - |\text{full}|)^2}
\]

where \(N\) is the number of data points, \(\text{full}\) is the fully sampled data set, and \(\text{data}\) is the under-sampled or MaxEnt reconstructed data set. Error bars are not shown because the standard deviations were three to four orders of magnitude smaller than the mean RMSEs and did not vary noticeably over percent under-sampling or SNR. As can be seen, the RMSE of the under-sampled data set increases as the percent of sampled data decreases but does not vary considerably with SNR. The MaxEnt reconstructions decrease the RMSE significantly at each SNR and percent under-sampling compared to the under-sampled data set but begin to rise at very low SNR.

The right pane of Figure 3 shows the average NAA to Cho and Lac to Cr ratios versus SNR for the fully sampled, under-sampled, and MaxEnt reconstructed data sets at 80%, 60%, 40%, and 20% under-sampling. The ratios were calculated from the peak integrated
areas of each metabolite at the ppm locations listed in Table 1 and then normalized by the fully sampled ratios at SNR=20. Each peak area was calculated only over the four spatially distributed voxels for each metabolite. Therefore, the NAA peak area was calculated over voxels (2,2), (2,3), (3,2) and (3,3) as shown in Figure 1, for example. This ensured that any spectral-spatial leakage caused by the under-sampling into adjacent voxels would modify the peak area for each metabolite and change the metabolite ratios as their peaks aliased along the $k_y$-$t_1$ plane.

The NAA/Cho and Lac/Cr ratios for the under-sampled data set shown in the bottom of the right pane in Figure 3 differ noticeably from the fully sampled ratios at each SNR. The NAA/Cho ratios are 1-8% larger than the fully sampled ratios and increase as the percent under-sampling decreases. The Lac/Cr ratios decrease as less data is sampled and are up to 9% lower than the fully sampled ratios for sample rates of 80, 60, and 40%, and 17% lower for the 20% sample rate. Both the NAA/Cho and Lac/Cr ratios show significant deviations in the metabolite ratios compared to the fully sampled true ratios; however, these deviations are limited to the percent under-sampling and are not largely affected by SNRs down to 2.

The MaxEnt reconstruction ratios for NAA/Cho and Lac/Cr shown in the top of the right pane in Figure 3 show much better agreement with the fully sampled ratios at each SNR and percent under-sampling than the under-sampled data. The NAA/Cho ratios are within 1% of the fully sampled ratios at SNRs greater than 6 but do deviate as high as 3% for lower SNRs. The Lac/Cr ratios are also improved across all SNRs to be within 5% of the fully sampled ratios for sample rates of 80, 60, and 40% and 12% for the 20% sample rate; however, unlike the NAA/Cho ratios, the Lac/Cr ratios were affected by decreasing SNR and improvements over the under-sampled data set are less significant at SNRs less than 6.

**Retrospective Under-sampling of 4D EP-COSI**

The results from retrospectively under-sampling to 25% and then MaxEnt reconstructing a fully sampled 4D EP-COSI scan of healthy breast in vivo are shown in Figure 4. The
mask used to under-sample the $k_y$-$t_1$ plane is shown on the far left along with the signal envelopes for each dimension and was generated using the 2D Poisson-gap method described earlier in this paper. The central point in $k_y$ was sampled for each $t_1$ increment and the 1st $t_1$ point was sampled at each $k_y$. [35] [17]. Because it is a 4D data set where each voxel is a 2D spectrum, space limitations prevent us from showing every voxel in the 16x16 FOV. Therefore, only a selected 2D COSY spectrum from the data set as well as the spatial distribution of specific spectral peaks is shown.

Figure 4 A1 shows a 2D COSY spectrum extracted from the fully sampled 4D EP-COSI scan of healthy breast in vivo. It was taken from the fatty breast region highlighted in Figure 4 A2. It clearly shows the lipid diagonal peaks: olefinic fat (UFD), methyl fat (FMETD), and fat (FAT/FAT2/FAT3) and the cross peaks: unsaturated fatty acid right (UFR), unsaturated fatty acid left (UFL), and triglyceryl fat (TGFR) [7]. The spatial distribution of the UFL and UFR cross-peaks from the fully sampled 4D EP-COSI scan is shown in Figure 4 A2 overlaid on the anatomical MRI.

Figure 4 B1 shows the same 2D COSY spectrum as Figure 4 A1 after 25% under-sampling has been applied to the $k_y$-$t_1$ plane using the mask shown at left. The spatial-spectral incoherent artifacts from the under-sampling manifest as smeared peaks along $t_1$, which is illustrated by the collapse of the peaks in the 1D projection along the $F_1$ dimension on the right. The aliasing of the large diagonal fat peaks around ($F_2$=2ppm, $F_1$=2ppm) obscure the much smaller UFL and UFR cross-peaks around ($F_2$=2.1ppm, $F_1$=5.4ppm). Figure 4 B2 shows the spatial distribution of the UFL and UFR cross-peaks and how spatial artifacts from under-sampling $k_y$-$t_1$ manifest as errant peaks in adjacent voxels when compared to the distribution of fully sampled peaks in Figure 4 A2.

Figure 4 C1 shows the 2D COSY spectrum from Figure 4 B1 after MaxEnt reconstruction has been applied. The majority of spectral artifacts seen in Figure 4 B1 have been removed and the metabolite peaks have been rebuilt in the 1D projection on the right of the figure. The aliasing caused by the diagonal fat peaks around ($F_2$=2ppm, $F_1$=2ppm) has been removed and the UFL and UFR cross-peaks around ($F_2$=2.1ppm,
$F_1 = 5.4$ ppm) are fully resolved. When compared to the fully sampled spectrum in Figure 4 A1, the MaxEnt reconstructed spectrum has equivalent spectral peaks, and all of the significant diagonals and cross-peaks are fully resolved. The errant peaks in the spatial distribution in Figure 4 B2 caused by the spatial under-sampling have been removed in Figure 4 C2.

The range of values for the $(UFL+UFR)/(FAT3+FAT2)$ integrated peak area ratio and average error from the fully sampled ratios for all 24 voxels in the localized VOI is shown in Table 2 for the under-sampled and MaxEnt reconstructed data sets. As can be seen, under-sampling caused the ratios to vary considerably from the fully sampled ratios with a high mean ratio error. The MaxEnt reconstructed ratios show much better agreement with the fully sampled ratios and have a much smaller mean ratio error.

**Prospectively Undersampled 4D EP-JRESI**

The results from the prospectively under-sampled 4D EP-JRESI scan of a human prostate with malignant lesions in the left and right base are shown in Figures 5 and 6. The 25% under-sampling mask with signal envelopes along the $k_y$ and $t_1$ dimensions is shown on the left of Figure 5. Select 2D JPRESS spectra from the 4D EP-JRESI data set are shown in Figure 6 with the spatial distribution of specific spectral peaks in Figure 5.

Figure 5 shows the spatial distributions of the Cit peaks highlighted in Figure 6 for the under-sampled and MaxEnt reconstructed data sets; the purple highlighted voxel corresponds to the location of the extracted 2D J-resolved spectrum in Figure 6A, the red corresponds to Figure 6B, and the blue corresponds to Figure 6C. Because of the sensitivity profile of the endorectal coil, the SNR of voxels close to the rectal wall is higher than the SNR of voxels further away [42]. The MaxEnt reconstruction in Figure 5B shows the SNR of the Cit peaks in healthy tissue decreasing further from the rectal wall, as expected. However, the spatial distribution of the under-sampled data set in Figure 5A shows aliased Cit peaks near the apex of the prostate as well as high SNR peaks within the rectum that should not be there. The spatial distribution of Cit in the
under-sampled data is noisier in general and shows significant spectral-spatial artifacts when compared to the MaxEnt reconstruction.

The top of Figure 6 shows results for the 25% under-sampled 4D EP-JRESI scan and the bottom shows results for the MaxEnt reconstruction of that scan. Because the scan was prospectively undersampled, there is no fully sampled data for comparison.

Figure 6B shows 2D J-resolved spectra extracted from the 4D NUS EP-JRESI data sets that are highlighted in red in Figure 5. As shown in the under-sampled spectrum in the top of the figure, the citrate (Cit) diagonal and its cross peaks around $F_2=2.6$ppm are not easily resolved. There is significant aliasing of the Cit peaks along $t_1$ as well as the fat peaks around $F_2=1.7$ppm. The bottom of the figure shows the same 2D J-resolved spectrum after MaxEnt reconstruction of the 4D NUS EP-JRESI data set. The Cit diagonal and cross peaks are now fully resolved with no aliasing along $t_1$ as well as the fat peaks around $F_2=1.7$ppm.

A high Cit to Cho ratio is normal in healthy prostate tissue, but a low Cit to Cho ratio is indicative of pathology [43]. Therefore, Figures 6A and 6C show 2D J-resolved spectra from the 4D EP-JRESI data set from tumors in the left and right base of the prostate that are highlighted in purple and blue in Figure 5, so they should have decreased Cit and elevated Cho. The presence of tumors was verified by biopsy with Gleason scores of 3+3 and prostatic specific antigen of 8. The under-sampled spectra in the top of Figure 6C show Cho peaks at $F_2=3.2$ppm that are highly aliased along $t_1$ and Figure 6A shows aliased Cit at $F_2=2.6$ppm from adjacent voxels. The MaxEnt reconstructions are shown in the bottom of the figure for the same 2D J-resolved spectra. The Cho peaks are no longer aliased along $t_1$ and the errant peaks have been removed.

**DISCUSSION**

This work addresses the long scan time requirements of 4D MRSI, which is one of the main issues preventing its widespread adoption. Non-uniform under-sampling was used to accelerate the 4D data acquisition and MaxEnt non-linear reconstruction was used to
rebuild the missing samples. This technique resulted in minimal losses in spatial-spectral resolution while reducing scan times considerably.

The simulated 4D EP-COSI data set results shown in Figure 3 demonstrate how the MaxEnt reconstruction performs at different levels of noise and percent under-sampling. The MaxEnt reconstruction decreases the RMSE significantly at each SNR and percent under-sampling compared to the under-sampled data set but begins to rise at very low SNR, indicating the inability of MaxEnt to fully reconstruct the data set when features and sampling artifacts are obscured by high levels of noise. The data fidelity constraint in equation (3) determines how closely the MaxEnt reconstructed points must be to the sampled points to within the standard deviation of noise, which increases as the noise floor increases. This increase in the noise floor effectively “loosens” the fidelity constraint, which allows the reconstructed points to vary from their sampled counterparts and increases the entropy of the reconstructed spectrum. As the entropy increases, so too do the non-linearity and RMSE of the reconstruction with a loose fidelity constraint [44]. The data fidelity constraint can be “tightened” beyond the standard deviation of the noise in an effort to reduce the RMSE and reconstruction non-linearity but this prevents MaxEnt from completely removing the spatial-spectral under-sampling artifacts close to the noise floor, which could potentially obscure small features [27].

The results from Figure 3 show the NAA/Cho ratios are consistent with the fully sampled data set ratios and offer significant improvement over the under-sampled data, even without using methods to reduce the reconstruction non-linearity [45]; however, the Lac/Cr ratios deviate slightly from the fully sampled ratios at higher percent under-sampling and significantly at 20% under-sampling. Upon further investigation, it was found that each metabolite area used to calculate the NAA/Cho and Lac/Cr ratios, with the exception of Lac, was stable and consistent with the fully sampled areas over SNR and percent sampling; however, the areas calculated for Lac were consistently low across all SNRs for each percent under-sampling. This was caused by a mismatch between the 2D Poisson-gap sample masks that were sine-bell modulated along $t_1$ and the filtered signal envelope of Lac, which was thought to be a sine-bell but upon further investigation
it was found to be a shifted sine-bell. This shift in the Lac signal envelope meant the mask failed to sufficiently sample the high SNR Lac points along $t_1$ for each percent under-sampling and prevented the full Lac peak area from being reconstructed by MaxEnt [45] [35] [34]. This discrepancy emphasizes the importance of the sample mask to the SNR of each reconstructed metabolite for 4D EP-COSI data and the dependence on the shape of standard filters applied prior to reconstruction [46]. For each metabolite in EP-JRESI data, however, the maximum SNR point along $t_1$ is always at $t_1=1$ and is not apodized by spectral filters prior to reconstruction; therefore, an exponentially damped sample mask that only considers $T_2$ signal losses in the time domain can be used to sample the high SNR points for each metabolite without the need to consider coherence transfer signal envelopes.

The Poisson-gap sampling masks used in these experiments were generated by a random Poisson distribution, which injects a degree of uncertainty into the MaxEnt reconstruction. It has proven to be a robust technique that generates masks with desirable PSFs, as shown in Figure 2, and the RMSEs of reconstructed data sets using different Poisson-gap sample masks are stable [32]. Previous attempts by our group to use deterministic masks that were not randomly generated were difficult to optimize and suffered from coherent aliasing which cannot be removed by MaxEnt reconstruction [47] [35]. Recent work in under-sampled multidimensional NMR data sets using deterministic sample masks has shown promise and could be adapted to 4D MRSI [48]. However, these sample masks are modulated by an exponential decay function, which is suitable for EP-JRESI data, but would need to be modified to use a sine-bell function for EP-COSI data. Poisson-gap sampling has the advantage that it is easily modified to accommodate different modulation functions and can therefore be used for many different scan types. It is unknown whether these new deterministic methods are as accommodating as Poisson-gap in that regard.

The healthy human breast results in Figure 4 show that under-sampling and MaxEnt reconstruction work well for filtered in vivo EP-COSI scans with a sine-bell signal envelope. The SNR-loss caused by $T_2$ decay that was not present in the simulated 4D EP-
COSI data set did not reduce the efficacy of the reconstruction. There was still sufficient SNR in the time domain to reconstruct the diagonal and cross peaks in the spectral domain. The improvement in the ratios of certain lipid peaks shown in Table 2 indicates the signal from the large diagonal fat peaks that had been aliased over the $k_y-t_1$ plane by the under-sampling was returned to the diagonal peaks by MaxEnt reconstruction and surfaced the smaller UFL and UFR cross peaks that had been obscured. The NUS data set in Figures 4B1 and 4B2 show artifacts spread along $t_1$ and $k_y$ as well as reduced spectral resolution along $t_1$ caused by the broad NUS PSF. The homogenous nature of fatty breast spectra made it difficult to determine from the figure if the spatial resolution along $k_y$ decreased as a result of the NUS PSF as expected. However, it is clear the effects of the NUS PSF along $t_1$ were removed by MaxEnt in Figure 4C1 and the errant spectral peaks in the spatial distribution were removed in Figure 4C2, suggesting the spatial PSF along $k_y$ was improved. Any spectral bleed from the spatial PSF of the EP-COSI pulse sequence along $k_x$ was orthogonal to the effect of the NUS PSF along $k_y$ and was not affected by the MaxEnt reconstruction.

Under-sampling of 4D MRSI data can be implemented in the pulse sequence on the scanner and successfully reconstructed using MaxEnt as shown in Figures 5 and 6 for the prospectively under-sampled EP-JRESI prostate data. Spatial-spectral artifacts caused by the broad NUS PSF were removed by MaxEnt to reveal spectra that were spatially localized in $k_y$ properly and lacked significant spectral aliasing in $t_1$. The Cit multiplet in the MaxEnt reconstructed data set showed a nicely J-resolved AB-type structure that had been aliased by convolution with the NUS PSF in the under-sampled data. No comparisons can be made to the fully sampled data set as was done for the retrospective EP-COSI data set; however, the mean (Cr+Cho+Spm)/Cit ratio for the MaxEnt reconstructed 2D JPRESS voxel shown in Figure 6B and the adjacent four voxels within the prostate, excluding the tumors in the left and right base, is 0.547 with a coefficient of variation of 21.2% which is consistent with reported literature [43] [49].

This work has demonstrated the potential for MaxEnt to accelerate the acquisition of 4D MRSI data in vivo into clinically acceptable scan times. However, because of its
similarity to CS reconstruction, quantitative comparisons between these techniques need to be made in order to determine the superior method for reconstructing 4D MRSI data and will be a topic of future research. Additionally, determining whether Poisson-gap or deterministic sample masks should be used and optimizing the modulation functions for 4D EP-JRESI and EP-COSI data sets and different spectral filters will be addressed in future papers.

CONCLUSIONS

We have shown that MaxEnt can successfully reconstruct under-sampled 4D MRSI data by reconstructing a mixed domain spectral-spatial plane. Simulated 4D MRSI data provided a quantitative characterization of the MaxEnt reconstruction at different percent under-sampling and SNR and it was shown that in vivo 4D EP-COSI and 4D EP-JRESI scans could be reconstructed. This acceleration translates into a clinically viable 5 minute EP-COSI breast scan and 6 minute EP-JRESI prostate scan.

ACKNOWLEDGEMENTS

Authors acknowledge the scientific support of Dr. Rajakumar Nagarajan for recording the in vivo NUS EP-JRESI data. This work was supported by: 1) a National Institute of Health (NIH) training grant #5T15 LM07356 (BLB), 2) an IDEA Expansion grant from the US Army Department of Defense (DOD) Breast Cancer Research Program (BCRP)#W81XWH-10-1-0743 (MAT), and 3) an IDEA grant from the US Army DOD Prostate Cancer Research Program (PCRP)#W81XWH-11-1-0248 (MAT).
References


23 Herman M, Strohmer T. Compressed sensing radar. In: 2008 IEEE Radar Conference (RADAR); 2008; Rome, Italy.


Table 1: PPM locations for metabolite peaks used in ratio calculations

<table>
<thead>
<tr>
<th></th>
<th>NAA</th>
<th>Cho</th>
<th>Cr</th>
<th>Lac</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diagonals (ppm)</td>
<td>(2.0,2.0),</td>
<td>(3.2,3.2),</td>
<td>(3.1,3.1),</td>
<td>(1.3,1.3),</td>
</tr>
<tr>
<td></td>
<td>(2.5,2.5),</td>
<td>(3.5,3.5),</td>
<td>(3.9,3.9),</td>
<td>(4.1,4.1)</td>
</tr>
<tr>
<td></td>
<td>(4.3,4.3)</td>
<td>(4.0,4.0)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cross Peaks (ppm)</td>
<td>(2.5,4.3),</td>
<td>(3.5,4.0),</td>
<td>N/A</td>
<td>(1.3,4.1),</td>
</tr>
<tr>
<td></td>
<td>(4.3,2.5)</td>
<td>(4.0,3.5)</td>
<td></td>
<td>(4.1,1.3)</td>
</tr>
</tbody>
</table>

Table 2: Range and mean error of (UFL+UFR)/(FAT3+FAT2) integrated peak area ratios for fully sampled, NUS, and MaxEnt reconstruction.

<table>
<thead>
<tr>
<th></th>
<th>Ratio Range</th>
<th>Mean Ratio Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full</td>
<td>0.1840 - 0.0634</td>
<td>N/A</td>
</tr>
<tr>
<td>MaxEnt</td>
<td>0.1604 - 0.0550</td>
<td>0.0188 ± 0.0048</td>
</tr>
<tr>
<td>NUS</td>
<td>0.4262 - 0.2180</td>
<td>0.1737 ± 0.0459</td>
</tr>
</tbody>
</table>
FIGURE LEGENDS

**Figure 1**: Simulated quad phantom illustration. (Top) Spatial distribution of an NAA, Cr, Cho, and Lac diagonal peak when fully sampled. (Bottom) Spatial distribution of the same NAA, Cr, Cho, and Lac diagonal peak as above with 25% under-sampling in the $k_y$-$t_1$ plane.

**Figure 2**: Poisson sample mask creation along the $k_y$-$t_1$ plane for EP-JRESI and EP-COSI. (Top) Sine and exponentially modulated values for $\lambda$. (Upper Middle) Poisson distributed values for each $\lambda$ indicating the sample gap between that point and other points in the sample distribution. (Lower Middle) Resulting 2D Poisson-gapped sample mask. (Bottom) Magnitude PSF of the 2D sample mask.

**Figure 3**: Metrics comparing the under-sampled, MaxEnt reconstruction, and fully sampled 4D EP-COSI simulated data. (Left) RMSE of under-sampled and MaxEnt reconstructed data versus SNR for 20, 40, 60, and 80% under-sampling. (Right) Comparison of the NAA/Cho and Cr/Lac ratios for fully sampled data to MaxEnt (Top) and under-sampled (bottom) reconstruction versus SNR for 20, 40, 60, and 80% under-sampling.

**Figure 4**: (Left) Under-sampling mask used to retrospectively under-sample the $k_y$-$t_1$ plane to 25% (Top) Selected 2D COSY spectrum from a 4D EP-COSI scan of healthy, fatty breast highlighted in the bottom images for A1) fully sampled B1) 25% under-sampling C1) MaxEnt reconstruction (Bottom) Spatial distribution of the olefinic cross-peaks of unsaturated fatty acids (UFL/UFR) highlighted in the 2D COSY spectrum for A2) fully sampled B2) 25% under-sampling C2) MaxEnt reconstruction.

**Figure 5**: Spatial distribution of the citrate peaks from a 4D EP-JRESI scan of malignant prostate highlighted in the 2D JPRESS spectra from Figure 6 for (A) 25% Under-sampled (B) MaxEnt reconstructed. (Left) Mask used to prospectively under-sample the $k_y$-$t_1$ plane to 25%.

**Figure 6**: Select 2D J-resolved spectra from a 4D EP-JRESI scan of malignant prostate for (top) prospectively 25% under-sampled and (bottom) MaxEnt reconstructed. A) Spectra highlighted in purple in Figure 5 showing elevated Choline and depressed Citrate in a malignant lesion. B) Spectra highlighted in red in Figure 5 showing elevated Citrate and normal Choline in healthy prostate. C) Spectra highlighted in blue in Figure 5 showing elevated Choline and depressed Citrate in a malignant lesion.
Figure 1: Simulated quad phantom illustration. (Top) Spatial distribution of an NAA, Cr, Cho, and Lac diagonal peak when fully sampled. (Bottom) Spatial distribution of the same NAA, Cr, Cho, and Lac diagonal peak as above with 25% under-sampling in the ky-t1 plane.
Figure 2: Poisson sample mask creation along the ky-t1 plane for EP-JRESI and EP-COSI. (Top) Sine and exponentially modulated values for λ. (Upper Middle) Poisson distributed values for each λ indicating the sample gap between that point and other points in the sample distribution. (Lower Middle) Resulting 2D Poisson-gapped sample mask. (Bottom) Magnitude PSF of the 2D sample mask.
Figure 3: Metrics comparing the under-sampled, MaxEnt reconstruction, and fully sampled 4D EP-COSI simulated data. (Left) RMSE of under-sampled and MaxEnt reconstructed data versus SNR for 20, 40, 60, and 80% under-sampling. (Right) Comparison of the NAA/Cho and Cr/Lac ratios for fully sampled data to MaxEnt (Top) and under-sampled (bottom) reconstruction versus SNR for 20, 40, 60, and 80% under-sampling.
Figure 4: (Left) Under-sampling mask used to retrospectively under-sample the ky-t1 plane to 25% (Top) Selected 2D COSY spectrum from a 4D EP-COSI scan of healthy, fatty breast highlighted in the bottom images for A1) fully sampled B1) 25% under-sampling C1) MaxEnt reconstruction (Bottom) Spatial distribution of the olefinic cross-peaks of unsaturated fatty acids (UFL/UFR) highlighted in the 2D COSY spectrum for A2) fully sampled B2) 25% under-sampling C2) MaxEnt reconstruction.
Figure 5: Spatial distribution of the citrate peaks from a 4D EP-JRESI scan of malignant prostate highlighted in the 2D JPRESS spectra from Figure 6 for (A) 25% Under-sampled (B) MaxEnt reconstructed. (Left) Mask used to prospectively under-sample the ky-t1 plane to 25%.
Figure 6: Select 2D J-resolved spectra from a 4D EP-JRESI scan of malignant prostate for (top) prospectively 25% under-sampled and (bottom) MaxEnt reconstructed. A) Spectra highlighted in purple in Figure 5 showing elevated Choline and depressed Citrate in a malignant lesion. B) Spectra highlighted in red in Figure 5 showing elevated Citrate and normal Choline in healthy prostate. C) Spectra highlighted in blue in Figure 5 showing elevated Choline and depressed Citrate in a malignant lesion.