REITERATIVE MINIMUM MEAN SQUARE ERROR ESTIMATOR

FOR DIRECTION OF ARRIVAL ESTIMATION

AND BIOMEDICAL FUNCTIONAL BRAIN IMAGING

BY

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ABSTRACT

Two novel approaches are developed for direction-of-arrival (DOA) estimation and functional brain imaging estimation, which are denoted as ReIterative Super-Resolution (RISR) and Source AFFine Image REconstruction (SAFFIRE), respectively. Both recursive approaches are based on a minimum mean-square error (MMSE) framework.

The RISR estimator recursively determines an optimal filter bank by updating an estimate of the spatial power distribution at each successive stage. Unlike previous non-parametric covariance-based approaches, which require numerous time snapshots of data, RISR is a parametric approach thus enabling operation on as few as one time snapshot, thereby yielding very high temporal resolution and robustness to the deleterious effects of temporal correlation. RISR has been found to resolve distinct spatial sources several times better than that afforded by the nominal array resolution even under conditions of temporally correlated sources and spatially colored noise.

The SAFFIRE algorithm localizes the underlying neural activity in the brain based on the response of a patient under sensory stimuli, such as an auditory tone. The estimator processes electroencephalography (EEG) or magnetoencephalography (MEG) data simulated for sensors outside the patient's head in a recursive manner converging closer to the true solution at each consecutive stage. The algorithm requires a minimal number of time samples to localize active neural sources, thereby enabling the observation of the neural activity as it progresses over time. SAFFIRE has been applied to simulated MEG data and has shown to achieve unprecedented spatial and temporal resolution. The estimation approach has also demonstrated the capability to precisely isolate the primary and secondary auditory cortex responses, a challenging problem in the brain MEG imaging community.

TABLE OF CONTENTS

ACKNOWLEDGEMENTS	II
ABSTRACT	III
TABLE OF CONTENTS	IV
TABLE OF FIGURES	VI
TABLE OF TABLES	IX
CHAPTER 1: INTRODUCTION	1
1.1 Overview	
1.2 DIRECTION OF ARRIVAL ESTIMATION	
1.3 NEURAL SOURCE LOCALIZATION	
1.4 Thesis Outline	4
CHAPTER 2: BACKGROUND	6
2.1 DIRECTION OF ARRIVAL ESTIMATION	
2.1.1 Physical Scenario and Signal Representation	
2.1.2 Signal Correlation and Spectral Super Resolution	
2.2 NEURAL SOURCE LOCALIZATION	
2.2.1 Neural Mechanism and Imaging Techniques	
2.2.2 The Leadfield Matrix	
2.2.3 Forward Signal Model and The Inverse Problem	
2.5 FILTER THEORY	
2.3.1 Multinle Signal Classification (MUSIC)	
2.3.3 Minimum Mean Sauare Error Estimators	
2.3.4 FOCal Underdetermined System Solution	
2.3.5 Discussion of Direction of Arrival Algorithms	
2.3.6 Discussion of Neural Localization Algorithms	
CHAPTER 3: RE-ITERATIVE SUPER-RESOLUTION	
3.1 Reiterative Minimum Mean Square Error Estimator	
3.2 RISR ALGORITHM	
3.3 SPECTRAL OVER-SAMPLING	
3.4 MULTIPLE-TIME PROCESSING	
3.4.1 Incoherent Integration	
3.4.2 Eigen Decomposition	
3.4.3 Additional Methods	
CHAPTER 4: SOURCE AFFINE IMAGE RECONSTRUCTION	44
4.1 SAFFIRE ALGORITHM	
4.1.1 Basic Algorithm	
4.1.2 AFFINE TRANSFORMATION OF SOLUTION SPACE	
4.1.5 Matched Filter Bank Initialization	
4.1.4 Energy Normalization 4.1.5 Noise Covariance Estimation	
4.1.6 Implementation	
1	

4.1.7 Reconstruction of Dipole Time Course	
4.2 Multiple-Time Processing	
4.3 MULTIPLE-STAGES FOR VOLUMETRIC CONSTRAINTS	55
CHAPTER 5: SIMULATION RESULTS AND DISCUSSION	
5.1 RISR	
5.1.1 Performance Metrics	
5.1.2 Basic Performance	
5.1.3 Temporal Robustness	
5.1.4 Spectral Super-Resolution	
5.1.5 Data Sample Support	
5.1.6 Calibration Error	
5.1.7 Colored Additive Noise	
5.1.8 Observation of Sparse Solution Convergence	
5.2 SAFFIRE	
5.2.1 Experimental Setup and MEG Configurations	
5.2.2 Single Dipole Activation	
5.2.3 Nearby Dipole Pair	89
5.2.4 Mirrored Dipole-Pair in Primary Auditory Regions	
5.2.5 Auditory Response	
5.2.6 Time-Course Reconstruction with Interference	
CHAPTER 6: CONCLUSIONS AND FUTURE WORK	104
6.1 Conclusions	
6.2 Future Work	

TABLE OF FIGURES

Figure 2-1. Physical setup of $M = 1$ source signal impringing on an ULA of N	
sensors	7
Figure 2-2. Structural composition of a neuron ^[19]	. 11
Figure 2-3. Biomagnetometer system with 151 channels during cortical MEG	
recording at the Hoglund Brain Imaging Center in KUMC	12
Figure 2-4. Magnetic field B of a current I	13
Figure 2-5. Sorted leadfield vector norms for dipoles in the brain sample space	.17
Figure 2-6. Eigen spectrum of the auto-correlation matrix of leadfield matrix B	17
Figure 2-7. Normalized inner-product of steering vectors	20
Figure 2-8. Block diagram of statistical filtering for estimation	24
Figure 3-1. Block diagram of the RISR operation	36
Figure 3-2. Block diagram of I2-RISR operation	40
Figure 5-1. Angular spectrum (in electrical degrees) of RISR with $K = 1$ snapshot,	
N = 10, $SNR = 35 dB$ and uncorrelated sources over the first 8 iterations	58
Figure 5-2. Angular spectrums of different algorithms with uncorrelated sources as	t
-90°, -72° and 0°, $K = 20$ samples, $SNR = 20 dB$. 59
Figure 5-3. Probability of separation of RISR, e-RISR and RISR with C as input	
versus SNR for uncorrelated sources, $K = 20$ samples	60
Figure 5-4. Angular root mean square error of RISR, e-RISR and RISR with C as	
input versus SNR for uncorrelated sources, $K = 20$ samples	61
Figure 5-5. Probability of separation of RISR, MUSIC and SS-MUSIC versus SNR	for
uncorrelated sources, $K = 20$ samples	62
Figure 5-6. Angular root mean square error of RISR, MUSIC and SS-MUSIC versi	IS
SNR for uncorrelated sources, $K = 20$ samples	63
Figure 5-7. Anecdotal results of RISR, MUSIC and SSMUSIC with sources at -90°,	
-72° (correlated with source at -90°) and 0°, $K = 20$ samples, $SNR = 35 dB$. 64
Figure 5-8. Probability of separation of RISR, MUSIC and SS-MUSIC versus SNR	for
correlated sources at -90°, -81° and an uncorrelated at 0°, $K = 20$ samples	. 65
Figure 5-9. Angular root mean square error of RISR, MUSIC and SS-MUSIC versu	IS
SNR for correlated source, $K = 20$ samples	. 66
Figure 5-10. Monte Carlo simulation result of POS for two uncorrelated signals ov	ver
a range of super-resolution factor L versus SNR for RISR and SS-MUSIC	
algorithms	. 67
Figure 5-11. Monte Carlo simulation result of ARMSE for two uncorrelated signal.	<i>S</i>
over a range of super-resolution factor L versus SNR for RISR and SS-MUSIC	7
algorithms	68
Figure 5-12. Probability of separation of RISR versus SNR for uncorrelated source	?S
at -90°, -81° and 0° and $K = 1, 2, 4, 8, 16$ and 500 snapshots	69
Figure 5-13. Probability of separation of SS-MUSIC versus SNR for uncorrelated	
sources at -90°, -81° and 0° and $K = 1, 2, 4, 8, 16$ and 500 snapshots	. 70

Figure 5-14. Effect of calibration error on RISR and SS-MUSIC at each antenna
element for three uncorrelated sources for amplitude error of 5% and phase
<i>error of 5%</i>
Figure 5-15. Probability of separation versus percentage calibration error in
amplitude and in phase for RISR and SS-MUSIC with three uncorrelated sources
at $SNR = 25 dB$ and $K = 40 snapshots$
Figure 5-16. Angular spectrum of estimated signal strengths for different algorithms
with 3 uncorrelated signals and addictive noise with mainlobe at around 45°74
Figure 5-17. Ten most dominant eigen values of matrix P at the 15th iteration for the
estimation of (a) a single signal source at -90° ; (b) two uncorrelated signals at
-90° and -45°; (c) three uncorrelated signals at -90°, -45° and 0°
Figure 5-18. Physical configuration of 150 MEG sensors relative to the brain77
Figure 5-19. Activation curve for single dipole activation
Figure 5-20. Mean Global Field Power and the sensors response of the simulated
<i>MEG</i> measurements corresponding to single dipole activation
Figure 5-21. The angled and front/top view of the location of the true active dipole
8981, which is 6.1 cm from the center of the brain, estimated by (a) SAFFIRE
using $K = 1$ snapshots; (b) SAFFIRE using $K = 4$ snapshots; (c) FOCUSS e; and
(d) MNE with a threshold of 70% of maximum estimated dipole strength
Figure 5-22. Dipole strength estimated by SAFFIRE, the neural activity index
resulted from LCMV algorithm, and the dipole strength estimated by MNE for
activated dipole 8981
Figure 5-23. True activation curve of dipole 8981 and reconstructed time course of
dipole activity by (a) SAFFIRE algorithm with $K = 4$ snapshots and (b) LCMV
with 400 snapshots to construct the data covariance matrix
Figure 5-24. The angled and front/top view of the location of the true active dipole
7952, which is 4.4 cm from the center of the brain, estimated by (a) SAFFIRE
using $K = 1$ snapshots; (b) SAFFIRE with $K = 4$ snapshots; and (c) FOCUSS 85
Figure 5-25. True activation curve of dipole 7952 and reconstructed time course of
dipole activity by (a) SAFFIRE algorithm with $K = 4$ snapshots and (b) LCMV
with 400 snapshots to construct the data covariance matrix
Figure 5-26. The angled and front/top view of the location of the true active dipole
3310, which is 2.0 cm from the center of the brain,
Figure 5-27. Dipole strength estimated by SAFFIRE on the left and the neural activity
index resulted from LCMV algorithm on the right for activated dipole 3310 88
Figure 5-28. True activation curve of dipole 3310 and reconstructed time course of
dipole activity by (a) SAFFIRE algorithm with $K = 4$ snapshots and (b) LCMV
with 400 snapshots to construct the data covariance matrix
Figure 5-29. Dipole strength estimated by SAFFIRE and the neural activity index
resulted from LCMV algorithm for the dipole pair 8570 and 8629
Figure 5-30. Angled and front 3D-view of dipoles 8570 and 8629, which are 1.09 cm
apart and about 5 cm from the center of the brain, and dipole estimated by
SAFFIRE using $K = 4$ snapshots

Figure 5-31. Reconstructed time course of dipole 8570 and 8629 using SAFFIRE and
the true dipole activation curve90
<i>Figure 5-32.</i> Angled and front 3D-view of the dipoles in the primary auditory regions
and dipole estimated by SAFFIRE using $K = 4$ snapshots
Figure 5-33. Reconstructed time course of dipole 1088 and 7952 in the auditory
region using SAFFIRE and the true dipole activation curve
Figure 5-34. Locations of the dipole pairs in primary cortex and secondary cortex for
the ASSR simulation
Figure 5-35. Activation curves of primary and secondary auditory regions with peaks
at $t = 1$ and 1.05 s, the mean global field power of the received signals (top
right) and the MEG sensors response of all 150 channels
Figure 5-36. An example of the first stage. Top: true dipoles responsible for ASSR
and the dipole locations estimated by SAFFIRE-2 with 3-of-5 detector in 3D plot
for data at around $t = 972$ ms (584th sample); bottom: MGFP plot from $t = 968$
ms to $t = 1.085$ with primary and secondary activation curves which are out of
scale. The black vertical line indicates $t = 972$ ms on the time axis
<i>Figure 5-37.</i> An example of the second stage. Top: true dipoles responsible for ASSR
and the dipole locations estimated by SAFFIRE-2 with 3-of-5 detector in 3D plot
for data at around $t = 1.025s$ (616th sample); bottom: MGFP plot from $t = 968$
ms to $t = 1.085$ with primary and secondary activation curves which are out of
scale. The black vertical line indicates $t = 1.025$ s on the time axis
Figure 5-38. An example of the third stage. Top: true dipoles responsible for ASSR
and the dipole locations estimated by SAFFIRE-2 with 3-of-5 detector in 3D
plot for data at around $t = 1.052s$ (632^{rd} sample); bottom: MGFP plot from $t =$
968 ms to $t = 1.085$ with primary and secondary activation curves which are out
of scale. The black vertical line indicates $t = 1.052$ s on the time axis
Figure 5-39. Activation curves of dipoles 1088 and 7952 in the primary cortex with
peak delay of 500 ms and the MGFP plot of the MEG sensor signal
Figure 5-40 Angled and front 3D-view of dipoles 1088 and 7952 in the primary
cortex with 500 ms peak delay and the dipole estimated by SAFFIRE. The
bottom plot is the reconstructed time course of the
<i>Figure 5-41.</i> Angled and front 3D-view of dipoles 1088 and 7952 in the primary
cortex with 500 ms peak delay and the dipoles estimated by SAFFIRE. The
bottom plots are the reconstructed time course of the dipoles 1088 and 7952
and the true activation curve

TABLE OF TABLES

Table 3-1. RISR and MMSE variables	
Table 5-1. Ideal time indices for first to third stages of ASSR estimation. The	bracket
value is the sample index corresponding to the time	
Table 5-2. Estimated time of ASSR dipole localization for the three processing	g
schemes	

CHAPTER 1: INTRODUCTION

1.1 Overview

Numerous filter techniques have been developed over the past few decades to solve a branch of problems consisting of a sensor array receiving signals from single or multiple sources transmitted through some medium. The goal of these techniques is to localize the signal source in either two or three dimensions by applying an adaptive filter to the signal received at the sensor array. The direction of arrival estimation problem addressed in this thesis falls directly under this category. Although the neural signal reconstruction problem addressed in the rest of this thesis comes from a very different area of signal processing application, it belongs to the aforementioned branch of problems because of the signal modeling of the system. The proposed solutions to the direction of arrival estimation and to the neural signal reconstruction were formulated based on the same mathematical framework, which was inspired from a radar adaptive pulse compression processing technique. Due to the degradation of most existing DOA estimation methods under correlated signals, the robustness to source correlation contributes to one of the most important advantages of the proposed approach. These two algorithms also achieve high spatial resolution compared to other approaches for the respective problems. Other applications of this general estimator framework include radar signal processing [1], telecommunications [2], etc.

The introduction and motivation for each of the two algorithms is discussed in the next two subsections.

1.2 DIRECTION OF ARRIVAL ESTIMATION

Direction of arrival (DOA) estimation has been an area of active research in the past decades with strong focuses on the applications in communications, radar and medical imaging [3]. The physical scenario for the estimation problem consists of narrowband radio frequency (RF) signals, which are transmitted by sources in the far field, impringing in the form of planewaves upon some sensors in a particular spatial arrangement. A common type of sensor configuration is the uniform linear array (ULA) antenna [4, 5], which is composed of equally spaced sensor elements. With the planewave assumption, the received signal at a sensor can be mathematically expressed as the summation of signal energy from all directions in space with phase delays across sensor elements.

Matched filtering, i.e. the conventional beamformer [4], and Multiple SIgnal Classification (MUSIC) [6, 7] are among the more commonly used signal processing approaches for this problem. However, the low resolution of matched filtering and the signal correlation intolerance of MUSIC are drawbacks that lead to further research on DOA estimation techniques. Some filtering methods such as spatiallysmoothed MUSIC (SS-MUSIC) [8] and the Least-Squares-based FOCal Underdetermined System Solver (FOCUSS) [9,10] are developed to fix the drawbacks of the previous approaches yet with the cost of lower spectral resolution, increased amount of prior knowledge, filter regularization, etc. These methods will be introduced in more details in the Chapter 2. Inspired by the work done for a radar adaptive pulse compression (APC) algorithm [1], the approach proposed in this study, denoted as Re-Iterative Super-Resolution (RISR), is a parametric estimator that iteratively estimates the spectral power by adaptively updating the filter bank at each stage without the need of prior knowledge. RISR tolerates correlated signals as well as colored noise, while also achieves angular super-resolution. This approach will be thoroughly discussed throughout Chapter 3.

1.3 NEURAL SOURCE LOCALIZATION

Electroencephalography (EEG) and magnetoencephalography (MEG) are imaging techniques used in clinical and research settings to measure the electrical and magnetic fields generated directly by the electrical activity of the brain. The main clinical application of MEG is functional brain imaging in which it allows non-invasive detection of neural activities, ensuring the safety of the patient. Brain imaging involves estimating the locations of brain activity based on the recordings of sensor elements surrounding the head. In essence, this localization problem is the extension of the direction of arrival estimation in which the location of a region of brain activity is estimated. Signal processing techniques are developed to determine the precise spatial location of underlying active neural sources [4]. It is done by processing measurements obtained by an array of MEG/EEG sensors outside the head, for which the electric and magnetic characteristics are modeled mathematically.

This estimation problem has been a topic of intense research in the past two decades and some examples of these signal processing techniques include the linearly constrained minimum variance beamformer (LCMV) [11,12] and the Focal Underdetermined System Solution (FOCUSS) [9,10]. Since LCMV assumes no temporal or spatial correlation between the neural source signals at different locations in the brain, it is prone to result in erroneous signal cancellation when such correlation is present [8, 13, 14]. The need for availability of relatively long stretches of data (to obtain good approximation of the data covariance matrix) in the case of LCMV and the required prior knowledge of interferer for multiple constrained minimum-variance beamformers with coherent source region suppression (MCMV-CSRS) [15], which is a modified version of LCMV, are the main drawbacks that limit the use of this class of algorithm. The initial estimates of FOCUSS algorithm tend to be biased towards locations close to the surface of the brain, which is a result of ill-conditioning of the transformation matrix in the forward model caused by the attenuations along the transmitted path of MEG signals. As a result, FOCUSS tends to return incorrect estimated locations of neural activity when the underlying neural source is deeper within the brain. The mathematic formulations and the characteristics of some existing approaches are included in the next chapter.

The approach presented in this paper, denoted as the Source AFFine Image Reconstruction (SAFFIRE) algorithm, resolves the drawbacks of most existing algorithms and possesses a number of advantages. It operates within an affinetransformed solution space to eliminate the depth bias in the solution due to neural signal attenuation with the source-to-sensor distance. A matched filter bank initialization is used to provide a low-resolution estimate in order to ensure the inclusion of the true solution. The ability to operate on as few as one time-sample allows very high temporal resolution compared to the previous approaches as well as temporal correlation robustness. It has been shown through simulations that SAFFIRE is also highly tolerant of spatial correlation of neural sources, a case in which LCMV fails. In simulations, this algorithm has successfully separated the simulated activity with the primary and secondary auditory cortex, which is a challenging problem in the MEG/EEG imaging community. The promising results of this algorithm obtained thus far should motivate further studies of its application in other branches of biomedical imaging problems.

1.4 THESIS OUTLINE

A presentation of several existing signal processing approaches and the classification of the estimation methods are included in Section 2.1 of Chapter 2. It is followed by some background material, such as definitions and the physical configuration of models, of the two problems addressed in this thesis. These sections are necessary since the proposed algorithms are application-specific and require more than just the understanding of the mathematical theories of the filters. The advantages and drawbacks of some existing algorithms are also discussed at the end of each section to compare the performance of RISR and SAFFIRE with other algorithms in their corresponding application area.

Although the direction of arrival estimation problem addressed by RISR is more general than the dipole localization by SAFFIRE, each of them is very complex in its own way, which leads to different implementations of the respective algorithm. Therefore, our two algorithms are introduced and discussed separately in individual chapters. Chapter 3 covers the basis and derivation of the RISR algorithm for direction of arrival estimation and its modification to enable spectral super-resolution and multiple-data-snapshots processing. The SAFFIRE algorithm is thoroughly described in Chapter 4 with the justification of some steps in the algorithm and how they dramatically change the performance of the estimator due to the ill-conditioning of the forward model transformation matrix. Multiple-time processing of SAFFIRE is introduced to allow higher signal-to-noise ratio in the estimation. The next section investigates the extension of the basic SAFFIRE algorithm into a multiple-stage processing scheme which produces volumetric constraints at each stage to confine the region of neural activity for the processing at the next stage.

The simulation results of both algorithms are presented in Chapter 5 along with the discussions and findings from each simulation case. Chapter 6 summarizes the work demonstrated in this thesis and gives insights on possible future research work pertaining to these estimation methods.

CHAPTER 2: BACKGROUND

2.1 Direction of Arrival Estimation

Determining the direction of which signal sources are transmitted can be achieved by sampling the spatio-temporal signal using a sensor array. The discretized signal is processed using some signal processing technique through which the information about the number of signals and/or their directions is extracted. Before introducing the signal processing approaches, it is essential to model the physical setup mathematically so that the quantity or arguments in the DOA approaches possess some physical representation.

2.1.1 PHYSICAL SCENARIO AND SIGNAL REPRESENTATION

The most typical category of signals is the narrowband signal, where the message signal of the source occupies a bandwidth considerably smaller or "narrower" than its carrier frequency. In our work, it is also assumed that the narrowband signal is transmitted in the far field to allow the plane wave approximation of the received signal at the sensors so that the curvature of the propagated wave can be ignored in our model. As a result of the narrowband assumption under the far field condition, there are two contributions to the difference of the received signal across sensors, which are the angular direction of the signals and phase change due to the carrier frequency.

The configuration of a 2-dimensional uniform linear array (ULA), as illustrated for the case with only a single source in Fig. 2-1, consists of N equally spaced sensors or antenna elements. Analogous to the Nyquist Sampling Theorem, the antenna elements are required to be separated by a distance d less than half the wavelength of the carrier signal to avoid ambiguity caused by spatial aliasing. An antenna element spacing of half the carrier wavelength will be used throughout our work presented in this thesis unless stated otherwise.



Figure 2-1. Physical setup of M = 1 source signal impringing on an ULA of N sensors

The signal received at the n^{th} sensor at time t_o can be represented by a summation of signal energy from all directions impringing on the sensor:

$$y_{n}(t_{o}) = \int_{-\frac{\pi}{2}}^{\frac{\pi}{2}} x(\phi, t_{o}) e^{j2\pi(n-1)(d/\lambda)\sin(\phi)} d\phi$$
(2.1)

where $x(\phi, t_o)$ is the amplitude of signal from angle ϕ at time t_o , d is the distance between two consecutive sensors and λ is the wavelength of the transmitted signal carrier frequency. The complex exponential contains the phase information of the n^{th} sensor as a function of the wavelength of carrier sinusoid signal λ and the arrival direction of the signal.

Assuming the sources are sparse, which means there are only a limited number of non-zero localized energy signals, the number of underlying signals is much less than N, and the noise is additive, the received signal at the n^{th} sensor at time index k in the presence of additive noise can be approximated in discrete form as:

$$y_{n}(k) = \sum_{i=1}^{M} x_{i}(k) e^{j2\pi(d/\lambda)\sin(\phi_{i})(n-1)} + \upsilon(k) \qquad n = 1, ..., N$$
$$= \sum_{i=1}^{M} x_{i}(k) e^{j\theta_{i}(n-1)} + \upsilon(k) \qquad (2.2)$$

where *M* is the number of distinct source signals arriving at the *n*th sensor, $\mathbf{x}_i(k)$ is the amplitude of the *i*th source signal at time index *k*, ϕ_i and θ_i are the direction of arrival and the electrical angle corresponding to the *i*th source, respectively, and $\mathbf{v}(k)$ is the additive noise. The (*n*–1) quantity in the complex exponential corresponds to the phase shift across different sensors. Since the distance between sensors *d* and the carrier frequency wavelength λ are fixed in the problem, these quantities are combined with the angular direction of the signal ϕ_i to form a new quantity called the electrical angle through the following equation:

$$\theta_i = 2\pi \frac{d}{\lambda} \sin(\phi_i) \tag{2.3}$$

The range of physical angle of source directions ϕ_i from $-\pi/2$ to $\pi/2$ translates non-linearly into an electrical angle range from $-\pi$ to π . The conversion to electrical angle simplifies the problem formulation so that some quantities can be pre-computed once the fixed parameters such as sensors spacing are known.

For simplicity, single snapshots are considered and the notation for time index k will be dropped for most of the remaining sections. A snapshot is referred as a simultaneous sampling of received signals across all array sensors. Notice also that we make no assumptions about the nature of the additive noise in the signal model.

To consider the received signals generated by M signal sources across N sensors, the expression in Equation (2.2) can be written more compactly in matrix form as

$$\mathbf{y} = \mathbf{S}\mathbf{x} + \mathbf{v} \tag{2.4}$$

where **y** is a $N \times 1$ vector of the received signal samples, **x** is a $M \times 1$ vector of source signal strength at a particular time, **v** is the $N \times 1$ additive noise vector and **S** is a $N \times M$ matrix consisting of *M* steering vectors of length *N* corresponding to *M* electrical angles given by

$$\mathbf{S} = \begin{bmatrix} 1 & 1 & \dots & 1 \\ e^{j\theta_1} & e^{j\theta_2} & \dots & e^{j\theta_M} \\ e^{j2\theta_1} & e^{j2\theta_2} & \dots & e^{j2\theta_M} \\ \vdots & \vdots & \ddots & \vdots \\ e^{j(N-1)\theta_1} & e^{j(N-1)\theta_2} & \dots & e^{j(N-1)\theta_M} \end{bmatrix}$$
(2.5)

In the signal model equation (2.4), the vector \mathbf{y} is the received signal represented as a linear combination of weighted steering vectors from a set of electrical angles uniformly sampled from $-\pi$ to π , i.e.

$$\theta_i = \frac{2\pi}{N} (i-1) - \pi \qquad i = 1, \dots, N$$
(2.6)

Applying Nyquist sampling theorem to spatial sampling, the nominal resolution of the electrical angle with *N* sensors is $2\pi/N$ radians. Super-resolution refers to the capability of distinguishing signal sources closer than $2\pi/N$ radians [16]. The construction of the original RISR algorithm uses sampling of $2\pi/N$ in electrical angle, **x** and **S** become a $N \times 1$ vector and a $N \times N$ matrix, respectively, while **y** remains the same size. The steering matrix S in this case has of the same form as a DFT matrix, which is full rank and invertible. An element in vector **x** is non-zero if the associated angle matches with any of the *M* directions of source signals. To expand RISR for super-separation of two signal sources, *M* would be a lot larger than *N*.

2.1.2 SIGNAL CORRELATION AND SPECTRAL SUPER RESOLUTION

Some DOA estimation algorithms, such as MUSIC, degrade significantly or fail when the source signals are correlated [8, 14]. There exist two kinds of source signal correlation: temporal and spatial. When two sources are close together so that their signals have very similar "signature" across the sensors, the sources are said to be spatially correlated. In mathematical terms, the magnitude of the inner product of the two equal-norm steering vectors corresponding to the two directions is very close to the norms of the individual steering vectors. In general, spatial correlation has indirect impacts on the spatial resolution of some estimation algorithms because the ability of resolving two sources depend greatly on how different the two sources signals "look" at the sensors. Matched filter is one of the algorithms that suffer from low spatial resolution and it will be discussed in detail in the next chapter.

Temporal correlation occurs when the knowledge of the message signal transmitted by one source provides some knowledge of another source message signal, regardless of the locations or directions of the two sources. A simple example of temporal correlation is when the two message signals from different directions have identical complex phases, in which case a peak will appear in the cross correlation between the two signals. In practice, this could occur in scenarios involving multipath propagation or smart jammers.

Temporal correlation of two sources directly influences the performance of algorithms based on eigen-methods [8, 14], which can be explained intuitively by considering the sum of two individual received data vectors, each corresponding to one of the two different signal directions. If the vectors are temporally correlated completely, for example, the phase change of one signal is identical to that of the other, then the data covariance matrix, calculated by averaging the outer products over time, contains only one dominant eigen vector since the signals cannot be discerned through the average.

2.2 NEURAL SOURCE LOCALIZATION

2.2.1 NEURAL MECHANISM AND IMAGING TECHNIQUES

Neurons are the basic units responsible for the processing and transmission of neural signals in the human brain. Neurons are made up of soma, which is the cell body containing a nucleus, axon, and dendrites, as shown in Figure 2-2. Typically, dendrites act as receivers of electrical synapses (or connections), which are electrical stimulation, transmitted by another neuron. The signal travels to the soma from which an output signal is then transmitted through the axon. During synaptic transmission, a spatial change of ion concentration in dendrites generates a current flow in the neuron. According to Maxwell's equations, electrical current induces electrical and magnetic fields with a pattern depending on the current distribution. Neuromagnetic fields are weak magnetic fields generated by tens of thousands of synchronously activated neurons within a small spatial extent. The field strengths can be detected using superconducting quantum interference device (SQUID) [17, 18] magnetometer outside of the head in a non-invasive manner as shown in Figure 2-3.



Figure 2-2. Structural composition of a neuron [19]



Figure 2-3. Biomagnetometer system with 151 channels during cortical MEG recording at the Hoglund Brain Imaging Center in KUMC

One of several ways to construct a model for the neural mechanism responsible for the MEG signal is called multiple dipole method. It divides the entire brain volume into small grids and approximates the localized current flow due to neural mechanism, also known as the primary current, in each of the grids as a current dipole [20]. Each dipole can be characterized by its location, orientation, and current strength. The brain and its surrounding tissues can be modeled in a first approximation as a spherically symmetric homogenous conductor. The magnetic field **B** pattern of a current stagement I, as shown in Figure 2-4, indicates that only current dipoles with a component tangential to the conductor surface can generate magnetic field to be detected by SQUID magnetometer. Therefore radial current dipole does not contribute to the MEG measurements [21], which is not a problem since most of the sensory cortical regions are located at fissures that guarantee tangential current dipole components.



Figure 2-4. Magnetic field B of a current I

While human brain structure has been studied and understood through the advancement in biological sciences in the past centuries, the bio-imaging technology enabled by the recent technological developments provides greater accuracy of the internal images of the brain without any surgical procedure [22]. Several important medical imaging techniques include computer-assisted x-ray tomography (CAT [23, 24, 25]), magnetic resonance imaging (MRI [23,24,26,27]), to reveal the anatomical structure of the brain with high-resolution yet static images. Relationships between functional purposes and the activation of certain regions in the brain can be investigated functional through neuroimaging methods such as single-photon-emission computed tomography (SPECT [23, 28]), positron-emission tomography (PET, [23, 29, 30]) and functional MRI (fMRI, [31]). These methods provide functional brain information at relatively low temporal resolution. Another drawback of SPECT and PET is that the patients are under radiation exposure, or strong static magnetic fields during data acquisition. fMRI has recently been developed to deliver real-time brain imaging, called real-time-fMRI (rtfMRI [32, 33]), but it measures neural activity indirectly based on the detection of blood oxygenation level change in the brain, whereas magnetoencephalography (MEG) is a direct measure of neural activity based on the electromagnetic fields generated by activated neurons.

As mentioned in the introduction section, electroencephalography (EEG) and magnetoencephalography (MEG) are the brain imaging techniques used to measure the electric and magnetic fields directly generated by neural activities of the brain. They are more superior to other imaging methods in terms of temporal resolution and based on the fact that they are completely non-invasive. EEG signals can be distorted by the uneven structure of the head as compared to the head model and therefore is less accurate than MEG signals in determining the spatial location of the current dipole. However, since the MEG signals attenuate at a greater degree than EEG signals with the source-to-sensor distance, MEG is more difficult to use for localizing dipoles deeper in the brain. The use of most MEG signal processing algorithms are limited by this problem, which will be discussed in more details in the next section.

The main application of MEG is functional brain imaging, which, through processing and analyzing sensor data, associates brain regions to particular functional purposes and the activation sequences. Since MEG has high temporal resolution and it is a direct measure of the neuron activity as opposed to the blood oxygenated level change around neurons, MEG can provide direct information on the dynamics of neural activity. MEG has applications in a broad range of areas ranging from cognitive neuroscience research to epilepsy and pre-surgical brain mapping.

2.2.2 THE LEADFIELD MATRIX

Under the conditions that the brain and its surrounding tissues (cerebro-spinal fluid, skull, skin) are modeled as a spherically symmetric homogenous volume conductor and localized primary current is approximated as current dipoles, the magnetic field at any location outside of the head as a function of the location, orientation and strength of a particular current dipole in the source space was derived by Ilmoniemi et al. [21] and by Sarvas [34] in accordance with Maxwell's equations as follows:

$$\mathbf{B}(\mathbf{r}) = \frac{\mu_0}{4\pi} \frac{F(\mathbf{r}, \mathbf{r}_Q) \mathbf{Q} \times \mathbf{r}_Q - (\mathbf{Q} \times \mathbf{r}_Q \cdot \mathbf{r}) \nabla F(\mathbf{r}, \mathbf{r}_Q)}{F(\mathbf{r}, \mathbf{r}_Q)^2}$$
where $F(\mathbf{r}, \mathbf{r}_Q) = a(ra + r^2 - \mathbf{r}_Q \cdot \mathbf{r})$

$$\nabla F(\mathbf{r}, \mathbf{r}_Q) = (r^{-1}a^2 + a^{-1}\mathbf{a} \cdot \mathbf{r} + 2a + 2r)\mathbf{r} - (a + 2r + a^{-1}\mathbf{a} \cdot \mathbf{r})\mathbf{r}_Q$$

$$\mathbf{a} = (\mathbf{r} - \mathbf{r}_Q), a = |\mathbf{a}|, and r = |\mathbf{r}|$$
(2.7)

 μ_0 is the permeability of free space. **r** and **r**_Q are position vectors of the current dipole in the source space and the MEG sensor, respectively, **Q** is a 3×1 vector representing the orientation and strength of a current dipole, **B** is a 1×3 magnetic field vector with each element corresponding to the magnetic field generated on one of the three orthogonal directions. The MEG sensors will measure the magnetic field component that is orthogonal on each sensor's area (a scalar value).

A few observations can be made from Equation (2.7). Since the radial component of \mathbf{Q} is parallel to \mathbf{r}_{Q} , the cross product between the vectors is zero, which means the radial component of any neural current dipole does not contribute to the magnetic field measurement $\mathbf{B}(\mathbf{r})$ outside the conductor. In other words, the MEG sensors are insensitive to radially oriented current dipoles. This is not a concern to the MEG application in brain imaging because, as mentioned in previous section, neural activities occur in the fissures of the brain, which are groves in the cortex where current dipoles have major tangential components, can be reliably recorded. Another observation is that current dipoles located at the center of conductor volume, i.e. $\mathbf{r}_{\mathbf{Q}} = 0$, $\mathbf{B}(\mathbf{r})$ is also zero. The closer a current dipole is to the center of the brain, the small its cross product is with \mathbf{Q} , and hence, the smaller $\mathbf{B}(\mathbf{r})$.

When multiple MEG sensors are present, $\mathbf{B}(\mathbf{r})$ due to a unit-strength current dipole can be expressed using a compact matrix representation, denoted as leadfield matrix. Let *N* be the number of MEG sensors, then each *N*×1 vector of a leadfield matrix corresponds to the magnetic field measurement of the N sensors generated by one of the three components (the ϕ , θ , and ρ in spherical coordinate system) of an unit current dipole. Therefore, a leadfield matrix due to a single unit-strength dipole is a $N \times 3$ matrix. For the MEG applications under the spherically symmetric volume conductor approximation, since the radial component of source dipoles does not produce magnetic field outside of the cortex, its associated leadfield vector can be eliminated to reduce the leadfield matrix to $N \times 2$ for a single dipole.

The source space, which can be either the whole brain or restricted to the cortical mantle depending on the application, can be divided into small grids each of which is represented by an unit current dipole. In this case, the leadfield matrix can be further extended to a collection of $\mathbf{B}(\mathbf{r})$ due to each of the current dipole locations. Let *M* be the number of source grids, the leadfield matrix is then a concatenation of the *M* $\mathbf{B}(\mathbf{r})$ sub-matrices, resulting a *N*×2*M* matrix. In order to sample the brain volume with good spatial resolution, *M* is typically a lot larger than *N*, which causes the leadfield matrix \mathbf{B} to be underdetermined.

Leadfield matrix is essentially a discrete representation of the magnetic field produced by neural activity at different locations throughout the cortex region. However, the use of this matrix requires consideration about the condition of the matrix. As mentioned in the previous section, magnetic fields attenuate rapidly with distance to the sensors and as a result, the **B**(**r**) associated with inner dipoles with small $|\mathbf{r}|$ are much smaller than that with the outer dipoles with large $|\mathbf{r}|$. This causes the norms of the column vectors in **B**(**r**) corresponding to inner dipoles to be a lot less than that to outer dipoles. Figure 2-5 below shows the range of norms the leadfield vectors take on where the average distance between grid points is 3mm.



Figure 2-5. Sorted leadfield vector norms for dipoles in the brain sample space

The leadfield matrix **B** has extreme norm discrepancy and very "correlated" column vectors has a wide eigen value distribution which, according to the definition of matrix condition number, means the matrix can be very ill-conditioned. Figure 2-6 shows the eigen spectrum of the extremely ill-conditioned leadfield matrix used for the computation in the research presented in this thesis. The wide eigenvalue spread is partly due to the underdetermined nature of the B matrix, which usually leads to non-unique solutions because the high-dimensional underlying signal \mathbf{x} is transformed by **B** into a space with much lower dimensions.



Figure 2-6. Eigen spectrum of the auto-correlation matrix of leadfield matrix B

The condition number of the leadfield matrix is the main reason for the performance degradation of some existing source localization algorithms when the underlying neural source is close to the center of the brain. A basin of attraction is a region in the solution space onto which if the initial solution falls, the algorithm would evolve to a particular solution point in that region. The basins of attraction for a system with an ill-conditioned matrix are very uneven and may strongly favor solutions to the source locations near the surface of the cortex. In other words, the ill-conditioning skews the source localization solution to one that is closer to the MEG sensors. For the case of inner dipole activation, the basin of attraction for the true solution may be too small for most initialization methods to yield an initial solution in. The way this problem can be fixed is explained in the later sections.

2.2.3 FORWARD SIGNAL MODEL AND THE INVERSE PROBLEM

With the knowledge of the approximated magnetic field generated by any current dipole in the source space in the form of a leadfield matrix, the MEG sensors response can be simulated for different types of neural activation, given the time-course of the neural activity. The process of simulating MEG sensor signals due to neural activity in the source space is called forward modeling. The leadfield matrix in this case is also known as the transformation matrix. Due to the superposition property of the magnetic fields, the total sensor response generated from multiple current dipoles can be represented as a linear sum of the sensor responses generated by each of the dipoles.

Let *N* and *M* be the number of sensors and dipole grid, respectively. Equation (2.8) below shows the forward model in matrix notation:

$$\mathbf{y} = \mathbf{B} \, \mathbf{x} + \mathbf{v} \tag{2.8}$$

where **B** is the $N \times 2M$ leadfield matrix, **x** is the $2M \times 1$ dipole component strength vector, **v** is the $N \times 1$ additive noise vector and **y** is the $N \times 1$ received signal vector. For example, if only the ϕ component of the *i*th and *j*th current dipole is activated with unit strength, the resulting sensors vector \mathbf{y} is the linear combination of the respective leadfield vectors plus additive noise.

The problem of determining the location and strength of some underlying current dipole responsible for MEG signals detected at the sensors is called the inverse problem. Since the SAFFIRE algorithm is based on a parametric MMSE framework [35], solving the inverse problem require the use of the forward model which will be shown in the next chapter. Due to the ill-condition of the leadfield matrix in the forward model, most parametric estimators suffer from biased dipole localization solution.

2.3 FILTER THEORY

2.3.1 MATCHED FILTER

Matched filter is the most straightforward way to estimate an underlying signal. It utilizes the fact that the inner-product of one vector with another yields the maximum value when the other vector is the Hermitian of the first. Consider a $N \times 1$ observed vector **y** that consists of the sum of a scalar multiple, x_i , of a $N \times 1$ vector **a**_i and a $N \times 1$ random additive noise vector **v** as shown in Equation (2.9).

$$\mathbf{y} = x_i \mathbf{a}_i + \mathbf{v} \tag{2.9}$$

A matched filter \mathbf{h} is one that maximizes the Signal-to-noise ratio (SNR), which is the expected value of the ratio of the signal and noise components with filter \mathbf{h} applied:

$$SNR = \frac{E\left\|\mathbf{h}^{H} x_{i} \mathbf{a}_{i}\right\|_{2}^{2}}{E\left\|\mathbf{h}^{H} \mathbf{v}\right\|_{2}^{2}} = \frac{E\left\|\mathbf{h}^{H} x_{i} \mathbf{a}_{i}\right\|_{2}^{2}}{E\left\{\mathbf{h}^{H} \mathbf{v} \mathbf{v}^{H} \mathbf{h}\right\}} = \frac{E\left\|\mathbf{h}^{H} x_{i} \mathbf{a}_{i}\right\|_{2}^{2}}{\mathbf{h}^{H} E\left\{\mathbf{v} \mathbf{v}^{H}\right\}\mathbf{h}}$$
(2.10)

where $\|\cdot\|_2^2$ is the ℓ^2 -norm and $E\{\cdot\}$ is the expectation of the quantity. The SNR is maximized when **h** is equal to the vector \mathbf{a}_i since the term in the numerator of Equation (2.10) is maximized when it contains an inner-product of \mathbf{a}_i with its

Hermitian. When the underlying linear model is extended into containing a transformation matrix \mathbf{A} and vector \mathbf{x} instead of a vector \mathbf{a}_i and scalar x_i , the matched filter becomes the Hermitian of the matrix \mathbf{A} .

Matched filter can be applied to any problem that contains linear underlying models. For the case of DOA model as defined by Equation (2.4), transformation matrix \mathbf{A} becomes steering matrix \mathbf{S} , whereas for the neural dipole localization problem, \mathbf{A} becomes the leadfield matrix \mathbf{B} .

Matched filter is the simplest to use and performs well when only a single source signal is present. However, it does not consider interference or colored noise, which might result in false signal detection. Matched filter might also fail if the columns of the transformation matrix A are linear dependent, which in that case the matched filter result might maximize SNR corresponding to the incorrect column vector. Another drawback is that the matched filter results in the least spatial resolution if the neighboring transformation space (column) vectors are highly correlated, meaning the inner-product of a vector **a** with other vectors spatially close **a** are almost as large as the inner product of the steering vector at 0 [rad] with other steering vectors are shown in Figure 2-7. The width of the mainlobe spans a wide range of angles which can make two spatially close-by signals un-resolvable.



Figure 2-7. Normalized inner-product of steering vectors

2.3.2 MULTIPLE SIGNAL CLASSIFICATION (MUSIC)

A class of filter for estimation based on the eigen representation of the signal space is called eigenspace method [7, 36]. They can be applied to problems where the observed signal can be written as a sum of complex sinusoids of different frequencies and additive white Gaussian noise (AWGN). MUltiple SIgnal Classification (MUSIC) [7], which is one of the eigen method algorithms, makes use of the orthogonality nature of the signal space and noise space for signal space projection.

Consider an observed signal vector **y** of length *M*, which consists of a sum of *P* sinusoids correspond to *P* different frequencies and AWGN, assuming M > P:

$$\mathbf{y} = \mathbf{x} + \mathbf{v} = \sum_{i=1}^{P} (\alpha_i \cdot \mathbf{s}_i) + \mathbf{v}$$
where $\mathbf{s}_i = \begin{bmatrix} 1 & e^{j\omega_i} & e^{j2\omega_i} & \cdots & e^{j(M-1)\omega_i} \end{bmatrix}^T$
(2.11)

 $[\cdot]^T$ denotes vector transpose. The $M \times M$ autocorrelation matrix of **y**, **R**_y, becomes the sum of the autocorrelation matrix of **x**, **R**_x, and the identity matrix scaled by the noise power σ_v^2 :

$$\mathbf{R}_{\mathbf{y}} = \mathbf{R}_{\mathbf{x}} + \sigma_{v}^{2} \mathbf{I} = \sum_{i=1}^{P} \alpha_{i}^{2} \mathbf{s}_{i} \cdot \mathbf{s}_{i}^{H} + \sigma_{v}^{2} \mathbf{I}$$
(2.12)

Performing the eigen decomposition on R_y:

$$\mathbf{R}_{\mathbf{y}} = \mathbf{V} \Lambda \mathbf{V}^{H}$$
where $\mathbf{V} = \begin{bmatrix} \mathbf{v}_{1} & \mathbf{v}_{2} & \cdots & \mathbf{v}_{M} \end{bmatrix}, \ \mathbf{v}_{i}^{H} \mathbf{v}_{j} = \begin{cases} 0 & i \neq j \\ 1 & i = j \end{cases}$

$$\Lambda = \begin{bmatrix} \lambda_{1} & & 0 \\ & \lambda_{2} & \\ & & \ddots & \\ 0 & & & \lambda_{M} \end{bmatrix}, \ \lambda_{1} \geq \lambda_{2} \geq \cdots \geq \lambda_{M}$$
(2.13)

 λ_i 's and v_i 's are called eigen values and eigen vectors, respectively. When noise is absent, $\mathbf{R}_{\mathbf{y}}$ in Equation (2.12) reduces to $\mathbf{R}_{\mathbf{x}}$. Since $\mathbf{R}_{\mathbf{x}}$ is of rank *P*, $\mathbf{R}_{\mathbf{y}}$ is also of rank *P* and thus λ_i are zeros for i = P+1, P+2, ..., M. This implies the *P* steering vectors in vector **x** span the same *P* dimensional subspace in \mathfrak{R}^{M} as the first *P* eigen vectors **v**_i, where i = 1, ..., P. This P-dimensional subspace is the signal-plus-noise subspace since noise correlation matrix is full-rank. The rest of the (M-P) eigen vectors, **v**_i, where i = P+1, ..., M, corresponding to the noise-only subspace spans a subspace orthogonal to the signal-plus-noise subspace.

MUSIC estimates the true signal frequencies by projecting the steering vectors at a particular frequency ω onto the noise-only subspace as follow:

$$P(\omega) = \frac{1}{\sum_{i=P+1}^{M} |\mathbf{s}^{H}(\omega) \mathbf{v}_{i}|^{2}}$$
(2.14)

If a signal is present at frequency ω_o , its steering vector $\mathbf{s}(\omega_o)$ is in the signal-plus-noise subspace and thus should be orthogonal to the noise-only subspace eigen vectors. Therefore the denominator in Equation (2.14) should be numerically close to zero, causing a peak at $P(\omega_o)$ in the $P(\omega)$ spectrum. Note that the peaks in the $P(\omega)$ spectrum indicate the true signal frequencies or directions but the values of $P(\omega)$ are irrelevant to the actual signal power.

MUSIC performs very well with good spectral resolution as long as the underlying signal model satisfies the assumed model in Equation (2.11). MUSIC also requires the prior knowledge of number of sinusoid signals P, which might not always be available or might be hard to attain. MUSIC fails when the number of receiver elements M is equal or less than P because $\mathbf{R}_{\mathbf{y}}$ eigen space would then be spanned entirely by signal steering vectors, resulting an empty noise-only subspace. MUSIC degrades when the additive noise is non-white and of which $\mathbf{R}_{\mathbf{v}}$ contains an eigen vector with significant noise power that belongs to the signal-plus-noise subspace.

In practice, the autocorrelation matrix $\mathbf{R}_{\mathbf{y}}$ is computed using the time-averaged cross-product of observed signal vectors \mathbf{y} . If two underlying signals are temporally correlated, they might be represented together as a subspace spanned by one eigenvector in $\mathbf{R}_{\mathbf{y}}$. This can lead to insufficient rank in $\mathbf{R}_{\mathbf{x}}$ and "bleach over" a

part of the signal-space into the noise-only space, causing the subspace projection to be incorrect.

A preprocessing scheme to resolve the signal-correlation intolerance of MUSIC algorithm is called spatial smoothing [8, 14]. Instead of time-averaging the outer products of observed vector \mathbf{y} to form a $M \times M$ autocovariance matrix $\mathbf{R}_{\mathbf{y}}$, each time-sample, or snapshot, of $M \times 1$ vector $\mathbf{y}(t)$ is sub-sectioned into M-L+1 overlapping vectors of length L. The k^{th} sub-section of $\mathbf{y}(t)$ is denoted as $\mathbf{y}_{\mathbf{k}}(t)$ where k = 1, ..., M-L+1:

$$\mathbf{y}_{\mathbf{k}}(t) = \begin{bmatrix} y(t,k) & y(t,k+1) & \cdots & y(t,k+L-1) \end{bmatrix}^T$$
 (2.15)

where y(t,k) is the k^{th} element of $\mathbf{y}(t)$. The spatially-smoothed autocorrelation matrix $\overline{\mathbf{R}}_{\mathbf{y}}(t)$ is defined as the "spatial" average a total of *M*-*L*+1 subsections of $\mathbf{y}_{\mathbf{k}}(t)$:

$$\overline{\mathbf{R}}_{\mathbf{y}}(t) = \frac{1}{M - L + 1} \sum_{k=1}^{M - L + 1} \mathbf{R}_{\mathbf{k}}(t)$$
where $\mathbf{R}_{\mathbf{k}}(t) = \mathbf{y}_{\mathbf{k}}(t) \cdot \mathbf{y}_{\mathbf{k}}^{H}(t)$
(2.16)

The term "spatial" in the averaging is due to the application of this scheme into direction of arrival linear arrays models. The overall $L \times L$ observed signal vector autocorrelation matrix $\overline{\mathbf{R}}_{y}$ is then just the time-average of all the $\overline{\mathbf{R}}_{y}(t)$'s. When two coherent signals are present, in which case \mathbf{R}_{x} in MUSIC algorithm becomes singular, spatial-smoothing essentially make the modified $\overline{\mathbf{R}}_{x}(t)$'s full-rank by using up some degrees of freedom in the original \mathfrak{R}^{M} space. As a result, signal-correlation intolerance of MUISC can be handled using this preprocessing scheme. The algorithm combining MUSIC with this preprocessing scheme is called Spatially-Smoothed MUSIC (SS-MUSIC).

Using spatial-smoothing scheme requires that the number of subsections M-L+1 must be greater or equal to the number of underlying sinusoids P in order for $\overline{\mathbf{R}}_{\mathbf{x}}(t)$'s to be non-singular. The size of subsections L must also be greater than P to allow noise-only subspace for the operation of MUSIC. Therefore the minimum

number of sensors M needed for SS-MUSIC is 2P, which is nearly twice as large as that required for the original MUSIC, indicating SS-MUSIC has less degrees of freedom and thus poorer spatial resolution than MUSIC for a fixed number of sensors.

2.3.3 MINIMUM MEAN SQUARE ERROR ESTIMATORS

The standard minimum mean square error (MMSE) estimator is a fundamental framework [35, 36] for several other filtering algorithms, including RISR. Its formulation minimizes the estimation error e(n) at time index n between the filter (estimator) output y(n) and the desired response d(n) as shown in Figure 2-8:



Figure 2-8. Block diagram of statistical filtering for estimation

When a finite-length transversal filter with N elements is applied to estimate the desired response d(n), the vectors in Figure 2-8 become:

$$\mathbf{x}(n) = \begin{bmatrix} x(n) & x(n-1) & \dots & x(n-(N-1)) \end{bmatrix}^{T}$$
$$\mathbf{w} = \begin{bmatrix} w_{0} & w_{1} & \dots & w_{N-1} \end{bmatrix}^{T}$$
$$y(n) = \mathbf{w}^{H} \mathbf{x}(n) = \sum_{k=0}^{N-1} w_{k}^{*} x(n-k)$$
(2.17)

The standard cost function *J* of the MMSE estimator is defined as the mean square error:

$$J = E\{\|e(n)\|^{2}\} = E\{e(n) e^{*}(n)\}$$

= $E\{(d(n) - \mathbf{w}^{H}\mathbf{x}(n))e^{*}(n)\}$ (2.18)

where $E\{\cdot\}$ is the expectation of the quantity. To obtain the optimal MMSE estimator \mathbf{w}_0 , the derivative of J with respect to w_k^* , k = 0, 1, ..., N-1, is taken to yield:

$$\frac{\partial J}{\partial w_k^*} = \frac{\partial}{\partial w_k^*} \left(E\left\{ \left(d(n) - \mathbf{w}^H \mathbf{x}(n) \right) e^*(n) \right\} \right) \qquad \text{for } k = 0, 1, ..., N - 1$$

= $E\left\{ x(n-k) e^*(n) \right\} = 0$ (2.19)

Expanding the error term $e^*(n)$ and $y^*(n)$ within and re-arranging terms to obtain a set of Wiener-Hopf equations:

$$E\left\{x(n-k)\left(d^{*}(n)-\sum_{i=0}^{N-1}w_{o,i}\ x^{*}(n-i)\right)\right\}=0$$
(2.20)

$$\sum_{i=0}^{N-1} w_{o,i} E\{x(n-k) \ x^*(n-i)\} = E\{x(n-k) \ d^*(n)\} \quad \text{for } k = 0, 1, ..., N-1$$
(2.21)

where $w_{o,i}$ is the *i*th element of \mathbf{w}_{o} filter. The optimal MMSE estimator formulation is obtained by writing the *N* Wiener-Hopf equations in matrix form as a function of auto-correlation matrix **R** and cross-correlation matrix **p**:

$$\mathbf{w}_{o} = \mathbf{R}^{-1}\mathbf{p}$$
where $\mathbf{R} = \begin{bmatrix} r(0) & r(1) & \cdots & r(N-1) \\ r^{*}(1) & r(0) & \cdots & r(N-2) \\ \vdots & \vdots & \ddots & \vdots \\ r^{*}(N-1) & r^{*}(N-2) & \cdots & r(0) \end{bmatrix}$

$$r(i-k) = E\{x(n-k) x^{*}(n-i)\} \qquad (2.22)$$

$$\mathbf{p} = E\{\mathbf{x}(n) d^{*}(n)\} = [p(0) \quad p(-1) \quad \dots \quad p(1-M)]^{T}$$

$$p(-k) = E\{x(n-k) d^{*}(n)\}$$

 $(\cdot)^*$ denotes the complex conjugate operator. The standard minimum mean square error (MMSE) estimator is also called Wiener filter. In order for the MMSE estimator to operate in an addition dimension in the observed data, the $N \times 1$ filter

vector \mathbf{w}_{0} can be extended to a $N \times M$ Wiener filter matrix \mathbf{W}_{0} and the formulation can be derived with matrix calculus to yield the following:

$$\mathbf{W}_{o} = \mathbf{R}^{-1}\mathbf{P}$$
where $\mathbf{R} = \begin{bmatrix} r(0) & r(1) & \cdots & r(N-1) \\ r^{*}(1) & r(0) & \cdots & r(N-2) \\ \vdots & \vdots & \ddots & \vdots \\ r^{*}(N-1) & r^{*}(N-2) & \cdots & r(0) \end{bmatrix}$

$$r(i-k) = E\{\mathbf{x}(n-k) \ x^{*}(n-i)\}$$

$$\mathbf{P} = E\{\mathbf{x}(n) \ \mathbf{d}^{H}(n)\} = [\mathbf{p}_{1} \quad \mathbf{p}_{2} \quad \dots \quad \mathbf{p}_{M-1}]^{T}$$

$$\mathbf{p}_{k} = E\{\mathbf{x}(n) \ d_{k}^{*}(n)\}$$
(2.23)

where $\mathbf{W}_{\mathbf{o}}$ is the MMSE estimator matrix, $d_k^*(n)$ is the k^{th} element of the $M \times 1$ desired response vector $\mathbf{d}(n)$, \mathbf{R} and \mathbf{P} are the auto-correlation and the cross-correlation matrix, respectively. The cost function J is then the square of 2-norm of the error vector $\mathbf{e}(n)$.

MMSE estimator formulation requires some prior knowledge of the auto-correlation of the received signal **R** and the cross correlation between the underlying and the received signal **p**, which might be difficult to compute or unattainable. Therefore, MMSE estimation is mainly used for inverse modeling or system identification problems. It is not intended for DOA estimation since the autocorrelation of $\mathbf{x}(n)$ in Equation (2.23) can only be approximated by timeaveraging observed samples, which could produce singular **R** matrix or poor statistical knowledge if insufficient data is collected. RISR formulation, as can be seen in the next chapter, is a modified version of the optimal Wiener filter except that prior statistical knowledge about the underlying signal is not required.

Linearly constrained minimum variance (LCMV) approach [11, 12], which is applied in a wide range of applications such as neural MEG beamforming, is covariance-based MMSE estimator. For the dipole localization problem, the LCMV filter minimizes the power of filter output **y** subject to the condition of unit response
at the each of the dipole location. With the filter output substituted with $\mathbf{y} = \mathbf{W}^T \mathbf{x}$, the mathematical problem statement is given by

$$\min_{W(q_o)} tr\{\mathbf{W}^{\mathsf{T}}(\mathbf{r})\mathbf{C}(\mathbf{x})\mathbf{W}(\mathbf{r})\} \text{ subject to } \mathbf{W}^{\mathsf{T}}(\mathbf{r})\mathbf{B}(\mathbf{r}) = \mathbf{I}$$
(2.24)

where **r** and **B**(**r**) are, as defined previously, the 3×1 dipole position vector and the corresponding leadfield vectors, respectively and the **C**(**x**) is the covariance matrix of the received data vector **x**. Using the Lagrange multipliers, the solution to the constrained optimization problem can be obtained as [36]

$$\mathbf{W}(\mathbf{r}) = \left\{ \mathbf{B}^{T}(\mathbf{r}) \mathbf{C}^{-1}(\mathbf{x}) \mathbf{B}(\mathbf{r}) \right\}^{-1} \mathbf{B}^{T}(\mathbf{r}) \mathbf{C}^{-1}(\mathbf{x})$$
(2.25)

LCMV indirectly handles the leadfield norm-biasing problem by normalizing its power estimates by the spatial noise spectrum. Therefore, the normalized estimated power of a particular dipole \mathbf{r} , denoted as the neural activity index (NAI), is calculated as

$$\mathbf{NAI}(\mathbf{r}) = \frac{tr\left\{ \mathbf{B}^{T}(\mathbf{r}) \mathbf{C}^{-1}(\mathbf{x}) \mathbf{B}(\mathbf{r}) \right\}^{-1}}{tr\left\{ \mathbf{B}^{T}(\mathbf{r}) \mathbf{Q}^{-1} \mathbf{B}(\mathbf{r}) \right\}^{-1}}$$
(2.26)

where \mathbf{Q} is the noise covariance matrix. The construction of the covariance matrix $\mathbf{C}(\mathbf{x})$ involves averaging the outer products of hundreds of data samples to acquire sufficient statistical information. Noise covariance matrix \mathbf{Q} is obtained in the same manner with pure-noise data samples.

2.3.4 FOCAL UNDERDETERMINED SYSTEM SOLUTION

An iterative nonparametric approach denoted as FOcal Underdetermined System Solution (FOCUSS) [9, 10] is a re-weighted minimum norm algorithm that determines the maximally sparse signal solution based on the minimization of the norm of the weighted estimates. Through iteratively updating the filter at each stage, the algorithm converges to a solution with localized energy from an initial lower resolution estimates. Although FOCUSS can be applied to non-linear problems, for the applications involved in this thesis assumes a linear transformation **A** of the unknown signal $\mathbf{x} \in C^{M}$ into its representation $\mathbf{y} \in C^{N}$ as

$$\mathbf{A}\mathbf{x} = \mathbf{b} \tag{2.27}$$

where **A** is the $N \times M$ transformation matrix, **x** and **b** are the $M \times 1$ unknown signal vector and the $N \times 1$ data vector, respectively. The weighted minimum norm solution for estimation of **x** is one that minimizes the ℓ^2 -norm of $\|\mathbf{W}^+\mathbf{x}\|$ subject to $\mathbf{AW}(\mathbf{W}^+\mathbf{x}) = \mathbf{b}$ where **W** is a $M \times M$ diagonal weighting matrix (not the filter matrix). The standard form of the solution is

$$\widetilde{\mathbf{x}} = \mathbf{W}(\mathbf{A}\mathbf{W})^{+}\mathbf{b} \tag{2.28}$$

where $\{\cdot\}^+$ denotes the Moore-Penrose (or pseudo-) inverse defined by $\mathbf{A}^+ = \mathbf{A}^H (\mathbf{A}\mathbf{A}^H)^{-1}$. The weighting matrix **W** is updated at each iteration stage with the signal estimates from the previous stage. The core components of FOCUSS algorithm are outlined below.

General FOCUSS algorithm

$$\mathbf{W}_{pk} = \mathbf{x}_{k-1}^{l} \circ \mathbf{I}_{M \times M}, \qquad l \in \mathbf{I}_{+}$$

$$\mathbf{q}_{k} = \left(\mathbf{A}\mathbf{W}_{ak}\mathbf{W}_{pk}\right)^{+}\mathbf{b} \qquad (2.29)$$

$$\widetilde{\mathbf{x}}_{k} = \mathbf{W}_{ak}\mathbf{W}_{pk}\mathbf{q}_{k}$$

I₊ denotes the set of all positive integers, \mathbf{W}_{pk} is the $M \times M$ diagonal weighting matrix containing the estimates \mathbf{x}_{k-1} from the previous stage raised to some power l and \mathbf{W}_{ak} is an additional $M \times M$ weighting matrix to allow flexibility for the algorithm for different applications. For example, leadfield-norm biasing for neural dipole localization can be compensated indirectly by assigning scaled leadfield norm differences to the diagonal elements of \mathbf{W}_{ak} .

Initialization is very important to a recursive algorithm in the sense that it initially narrows down the solution set from a larger set of possible solutions. FOCUSS utilizes the minimum norm estimate of the model in Equation (2.27) as the initialization, i.e. $\tilde{\mathbf{x}}(0) = \mathbf{A}^+ \mathbf{b}$. Depending on the application, standard minimum norm solution could yield an estimate from an incorrect basin of attraction in which the converged solution of FOCUSS resides. For example, a heavily ill-conditioned transformation matrix A can be characterized by very uneven basins of attraction, some of which contains a large solution set that can erroneously attract the minimum norm solution. As demonstrated in later chapters, for neural dipole localization this initialization method results in biased initial estimates in the wrong basin of attraction such that FOCUSS does not converge to the true solution.

Since noise is not built into the signal transformation formula, regularization to stabilize the matrix inverse in the filter formulation becomes an issue when implementing FOCUSS. The choices of regularization methods and how the optimal regularization parameter values are identified vary for different applications and are discussed in detail in the literature [9, 10].

2.3.5 DISCUSSION OF DIRECTION OF ARRIVAL ALGORITHMS

As mentioned in the introduction section, there exist classes of algorithms with different abilities to handle temporally correlated signals. The classical methods such as the MUSIC algorithm [7] break down when signals from different directions are partially or completely correlated in time. The main reason is that they require multiple data snapshots to construct or approximate the data covariance matrix of the signals, which might match with the outer product of steering vectors from another direction when the underlying signals are correlated in time.

Spatial-smoothing MUSIC [8] differs from MUSIC in terms of the construction covariance matrix from which the eigen vectors are decomposed. This modification to the algorithm increases its robustness towards correlated signal but its super-resolution performance degrades. Since MUSIC is only a detection algorithm, estimates of the underlying signals have to be obtained through a separate scheme; whereas RISR determines both signal strength and direction in the algorithm. SS-MUSIC requires prior knowledge of the number of signals to optimize

the size of subarrays to construct the spatially smoothed covariance matrix. The RISR estimator does not require any prior knowledge and is inherently tolerant of coherence signals.

FOCUSS is mathematically similar to RISR and it can operate on one snapshot as well. However as noted earlier, FOCUSS uses matrix inverse regularization to avoid ill-conditioning and the amount of regularization is a dual problem that must be solved, hence it is a less attractive approach than RISR. The performance comparisons of different algorithms on their robustness on temporal correlated signals are presented in the Chapter 5.

Since noise is not built into the signal transformation formula, regularization to stabilize the matrix inverse in the filter formulation becomes an issue when implementing FOCUSS. The choices of regularization methods and how the optimal regularization parameter values are identified vary for different application and are discussed in detail in the literature [9,10].

2.3.6 DISCUSSION OF NEURAL LOCALIZATION ALGORITHMS

Minimum-norm estimates (MNE) [37], FOCUSS [9, 10], MUSIC [7, 38] and LCMV algorithms have been applied to the MEG neural localization problem, though with a number of limitations associated with them. Signal correlation is known to cause performance deterioration of the MUSIC algorithm. Since the construction of the data covariance matrix C(x) for LCMV requires averaging the outer products of hundreds of data samples so as to acquire sufficient statistical information, LCMV provides accurate yet low resolution estimates of signal power. Another drawback of LCMV is the performance degradation due to temporal correlation of underlying dipole time courses, in which case LCMV may incorrectly localize active dipoles or reconstruct distorted time courses.

A number of modified versions of LCMV, such as LCMV with partial sensor coverage (LCMV-PSC) [39] and multiple constrained minimum variance beamformers with coherent source region suppression (MCMV-CSRS) [15, 40], have been formulated to solve the problem of interferer insusceptibility. However, the algorithms have a number of limitations, for example, LCMV-PSC reduces the degrees of freedom in the estimation by ignoring sensors sensitive to the interferers. In order to suppress coherent dipole activities, MCMV-CSRS requires prior knowledge of the location of the interferer, which might not always be available.

Instead of a data covariance matrix, a structured covariance matrix is built into SAFFIRE filter formulation, of which the diagonal elements of the matrix asymptotically approach the power of the dipoles in the sample space as the solution converges. As a result, SAFFIRE does not need to extract statistical prior knowledge from a large amount of data samples, enabling SAFFIRE to operate on as few as one data snapshot and achieve very finer temporal resolution.

The standard minimum norm estimate (MNE) method provides a solution that minimizes the power of the estimated dipole strengths and matches with the MEG sensors data [37]. The known limitations of this method include low spatial resolution due to high spatial correlations, and severe bias towards superficial dipoles induced by the large associated leadfield norms.

Despite that FOCUSS achieves finer resolution than LCMV and MNE, the bias compensation employed through the weight matrix at each iteration might provide proportionate weighting to the dipole estimates for some cases, therefore it requires consideration of the nature of the sensor environment and sampling space for construct an effective bias adjustment. Moreover, as mentioned previously, substantial amount of effort is involved in choosing regularization method and determining the optimal parameters to stabilize matrix inverse, which could varies depending on experimental setting.

Since the noise component is incorporated into the SAFFIRE algorithm through the noise covariance matrix, which can be approximated easily, there is no need for matrix inverse regularization. In order to avoid superficial dipole biasing, the filter operation of SAFFIRE is carried out in an affined-transformed space with energy normalization at each stage. This strategy effectively eliminates the leadfield norm biasing even for dipole located deep in the brain volume.

CHAPTER 3: RE-ITERATIVE SUPER-RESOLUTION

3.1 REITERATIVE MINIMUM MEAN SQUARE ERROR ESTIMATOR

As discussed in the previous chapter, the autocorrelation matrix \mathbf{R} and crosscorrelation matrix \mathbf{p} in the MMSE estimator formulation are unknown in advance for some applications. An adaptive approach denoted as Reiterative Minimum Mean Square Error (RMMSE) estimator, which was inspired by a radar pulse compression algorithm, is developed based on the MMSE framework [1, 35] to handle the lack of prior statistical knowledge. The fundamental differences between RMMSE and MMSE estimators are discussed later in this section.

Consider the following signal model, which has a similar form as Equation (2.4):

$$\mathbf{y}(n) = \mathbf{S}\mathbf{x}(n) + \mathbf{v}(n) \tag{3.1}$$

where $\mathbf{y}(\mathbf{n})$ is a $N \times 1$ observed signal vector, $\mathbf{x}(\mathbf{n})$ is a $N \times 1$ underlying signal vector, $\mathbf{v}(\mathbf{n})$ is the $N \times 1$ additive noise vector, and \mathbf{S} is a $N \times N$ transformation matrix. To relate RMMSE with the MMSE estimator framework in Chapter 2, their respective signal models should have equivalent variable notations, which are listed in Table 3-1.

Table 3-1. RISR and MMSE variables	
RMMSE	MMSE
x (n)	d (n)
y (n)	x (n)
$\mathbf{W}^{H}\mathbf{y}(n)$	y (n)

To derive RMMSE, Equation (2.23) can first be rewritten as

$$\mathbf{W} = \left(E\left\{ \mathbf{y}(n) \cdot \mathbf{y}^{H}(n) \right\} \right)^{-1} E\left\{ \mathbf{y}(n) \cdot \mathbf{x}^{H}(n) \right\}$$
(3.2)

Incorporating the signal model into the MMSE formulation by substituting in Equation (2.4) with a reasonable assumption that the additive noise is uncorrelated with the signal, the estimator becomes:

$$\mathbf{W} = \left(E \left\{ \left(\mathbf{S}\mathbf{x}(n) + \mathbf{v}(n) \right) \cdot \left(\mathbf{S}\mathbf{x}(n) + \mathbf{v}(n) \right)^{H} \right\} \right)^{-1} E \left\{ \left(\mathbf{S}\mathbf{x}(n) + \mathbf{v}(n) \right) \cdot \mathbf{x}^{H}(n) \right\}$$
$$= \left(E \left\{ \mathbf{S}\mathbf{x}(n)\mathbf{x}^{H}(n)\mathbf{S}^{H} + \mathbf{v}(n)\mathbf{v}^{H}(n) \right\} \right)^{-1} E \left\{ \mathbf{S}\mathbf{x}(n) \cdot \mathbf{x}(n)^{H} \right\}$$
(3.3)
$$= \left(\mathbf{S}\mathbf{R}_{\mathbf{x}}\mathbf{S}^{H} + \mathbf{R}_{\mathbf{v}} \right)^{-1} \left(\mathbf{S}\mathbf{R}_{\mathbf{x}} \right)$$

where $\mathbf{R}_{\mathbf{x}} = E\{\mathbf{x}(n) \cdot \mathbf{x}^{H}(n)\}$ is the *N*×*N* auto-correlation matrix of the underlying signal $\mathbf{x}(n)$, and $\mathbf{R}_{\mathbf{v}} = E\{\mathbf{v}(n) \cdot \mathbf{v}^{H}(n)\}$ is the *N*×*N* noise covariance matrix, which can be approximated by averaging across time the outer products of the pure-noise received signal.

As one would expect, adaptivity of RMMSE is achieved by re-iteratively updating the structured covariance matrix $\mathbf{R}_{\mathbf{x}}$ in the filter through which the signal estimates $\tilde{\mathbf{x}}(n)$ can be refined in successive stages. Assuming the elements in signal vector $\mathbf{x}(n)$ are statistically uncorrelated to allow an approximation of a diagonal structured covariance matrix, the final form of RMMSE at the k^{th} stage can be expressed as:

$$\mathbf{W}_{k} = \left(\mathbf{SR}_{\widetilde{\mathbf{x}}_{k}}\mathbf{S}^{H} + \mathbf{R}_{v}\right)^{-1}\left(\mathbf{SR}_{\widetilde{\mathbf{x}}_{k}}\right)$$
(3.4)
where
$$\mathbf{R}_{\widetilde{\mathbf{x}}_{k}} = \begin{bmatrix} \left|\widetilde{x}_{k-1,1}(n)\right|^{2} & 0 \\ & \left|\widetilde{x}_{k-1,2}(n)\right|^{2} & \\ 0 & \ddots & \\ 0 & & \left|\widetilde{x}_{k-1,N}(n)\right|^{2} \end{bmatrix}$$

 $\mathbf{R}_{\tilde{\mathbf{x}}_k}$ is the diagonal matrix of the estimated signal strengths from the previous iteration and $\tilde{x}_{k,i}(n)$ is the *i*th element of the estimated signal. Zero values on the off diagonal terms are the result of assuming that signals from different directions are temporally uncorrelated. The resulting estimate at the *k*th stage is then:

$$\widetilde{\mathbf{x}}_{k} = \mathbf{W}_{k}^{H} \mathbf{y}(n) = \mathbf{R}_{\widetilde{\mathbf{x}}_{k}} \mathbf{S}^{H} \left(\mathbf{S} \mathbf{R}_{\widetilde{\mathbf{x}}_{k}} \mathbf{S}^{H} + \mathbf{R}_{v} \right)^{-1} \mathbf{y}(n)$$
(3.5)

The above equation reveals some information about the matrix operation of the RMMSE estimator onto the observed signal. At the k^{th} stage of the RMMSE estimation Equation (3.5), the received signal $\mathbf{y}(n)$ is multiplied first by the matrix inverse, through which $\mathbf{R}_{\tilde{\mathbf{x}}_k}$ nulls out any active sources estimated from the previous re-iteration stage and \mathbf{R}_v indirectly cancel effects due to the additive noise \mathbf{v} in the signal model. The remaining data is then match-filtered by S^H to estimate the signal that had not been accounted for in the previous RMMSE stage. Finally, the signal energy that had been nulled out by the matrix inverse is multiplied back to the estimates through $\mathbf{R}_{\tilde{\mathbf{x}}_k}$ to produce the estimator at the current stage.

The matrix inverse in Equation (3.5) essentially suppresses signal components in $\mathbf{x}(n)$ according to their signal strengths. For example, if the *i*th entry along the diagonal in $\mathbf{R}_{\tilde{\mathbf{x}}_k}$ from the previous stage is close to zero, meaning there is only a small amount of signal energy in that component, the *i*th column in \mathbf{W}_k will have a small norm. Therefore multiplying that the Hermitian of the *i*th column of \mathbf{W}_k to the $\mathbf{y}(n)$ produces a small estimate of the *i*th component in $\tilde{\mathbf{x}}_k$ at that stage.

Different initialization methods can be implemented for the structured covariance matrix $\mathbf{R}_{\tilde{\mathbf{x}}_0}$, depending on the filter application. Those with more well-conditioned transformation matrices, such as the DOA estimation, are more flexible on the type of initialization; whereas applications with ill-posed signal model, neural dipole localization for example, requires caution when determining the initialization method for reasons that will be explained in later sections.

Despite of the fact that RMMSE approach is derived from the general MMSE formulation, the two filters have some fundamental differences that permit one of them to be used in certain problems where the other cannot. The MMSE formulation results in a set of estimated parameters of the signal model and it minimizes the mean square error between the estimates $\tilde{\mathbf{x}}(n)$ obtained through the MMSE filter and the actual signal strengths $\mathbf{x}(n)$. The formulation assumes the knowledge of auto-

correlation of the received signal strength and the cross-correlation between the received signal and the actual signal, therefore making it non-adaptive and non-iterative. On the contrary, RMMSE algorithm does not require any prior knowledge of the signal. The RMMSE filter is formed only as a mean to obtain the system coefficients, which are assumed to get closer to the truth signal at successive iteration as the solution converges. Given the knowledge from the previous re-iteration, the estimate at each re-iterative stage results in the smallest MSE possible. The self-refining of the solution due to adaptivity of the filter is the main advantage of RMMSE estimation algorithm over the original MMSE estimation.

3.2 RISR ALGORITHM

RMMSE approach can be directly applied to the direction of arrival problem by replacing the general signal model in Equation (2.4) with the uniform linear array signal model defined in Chapter 2. This application specific RMMSE algorithm is denoted as Re-Iterative Super-Resolution (RISR).

With the lack of prior knowledge on data statistics, at the initialization stage we assume equal weights on steering vectors of all angles by setting the diagonal elements of $\mathbf{R}_{\tilde{\mathbf{x}}_0}$ to unity. We also neglect the noise component in the signal model yielding an expression for the initialization stage MMSE filter equivalent to the Moore-Penrose pseudo-inverse of the steering vector matrix:

$$\mathbf{W}_0 = \left(\mathbf{S} \cdot \mathbf{S}^H\right)^{-1} \mathbf{S} \tag{3.6}$$

This filter is independent of the received data and therefore can be pre-computed. For the first re-iteration stage of the algorithm, the estimates from the initialization stage are used to update the diagonal terms of $\mathbf{R}_{\tilde{\mathbf{x}}_1}$ in the RISR estimator, which is then applied to the original received data y to form the set of first re-iteration stage estimate $\tilde{\mathbf{x}}_1(n)$:

$$\widetilde{\mathbf{x}}_{1}(n) = \mathbf{W}_{1}^{H} \mathbf{y}(n) = \mathbf{R}_{\widetilde{\mathbf{x}}_{1}} \mathbf{S}^{H} \left(\mathbf{S} \mathbf{R}_{\widetilde{\mathbf{x}}_{1}} \mathbf{S}^{H} + \mathbf{R}_{\mathbf{v}} \right)^{-1} \mathbf{y}(n)$$
where
$$\mathbf{R}_{\widetilde{\mathbf{x}}_{1}} = \begin{bmatrix} \left| \widetilde{x}_{0,1}(n) \right|^{2} & 0 \\ \left| \widetilde{x}_{0,2}(n) \right|^{2} & \\ 0 & \left| \widetilde{x}_{0,N}(n) \right|^{2} \end{bmatrix} = \left(\widetilde{\mathbf{x}}_{0}(n) \widetilde{\mathbf{x}}_{0}^{H}(n) \right) \circ I_{N \times N} \quad (3.7)$$

$$\widetilde{\mathbf{x}}_{0}(n) = \mathbf{W}_{0}^{H} \mathbf{y}(n) = \mathbf{S}^{H} \left(\mathbf{S} \cdot \mathbf{S}^{H} \right)^{-1} \mathbf{y}(n)$$

where $\tilde{x}_{0,k}(n)$ is the k^{th} element in the $\tilde{\mathbf{x}}_0(n)$. The time-index notation is dropped in this section for simplicity. The estimates $\tilde{\mathbf{x}}_1(n)$ are used to update the $\mathbf{R}_{\tilde{\mathbf{x}}_2}$ of the filter \mathbf{W}_2 for the second re-iteration stage in the same manner as Equation (3.7), which is mathematically equivalent to computing the Hadamard product (element-by-element multiplication, denoted as $\{\circ\}$) of the outer-product of $\tilde{\mathbf{x}}_1(n)$ and a $N \times N$ identity matrix. The corresponding filter \mathbf{W}_2 then multiplies the original received data $\mathbf{y}(n)$ to produce the signal strength estimate $\tilde{\mathbf{x}}_2(n)$. The process repeats until the estimate converges, which usually takes less than 10 re-iterations. Convergence of estimates is satisfied when the mean square difference between consecutive estimates is small than an acceptable value ε , i.e. $\|\tilde{\mathbf{x}}_k(n) - \tilde{\mathbf{x}}_{k+1}(n)\|^2 < \varepsilon$. Figure 3-1 demonstrates the general signal flow in the RISR algorithm.



Figure 3-1. Block diagram of the RISR operation

To summarize, the core portion of the RISR algorithm are listed below:

Implementation of the basic RISR algorithm

Initialization:

i) Compute estimate of noise covariance matrix over the interval of N_{noise} time samples that contains purely noise:

$$\hat{\mathbf{R}}_{\mathbf{v}} = \frac{1}{N_{noise}} \sum_{k=1}^{N_{noise}} \mathbf{v}(n) \, \mathbf{v}^{H}(n)$$

ii) Compute the initial estimate with the pseudo-inverse of steering-vector matrix S:

$$\widetilde{\mathbf{X}}_{0}(n) = \mathbf{W}_{0}^{H} \mathbf{y}(n) = \mathbf{S}^{H} \left(\mathbf{S} \, \mathbf{S}^{H} \right)^{-1} \mathbf{y}(n)$$

Recursive stages:

For k = 1, 2, ..., K iterations, compute:

1)
$$\mathbf{R}_{\tilde{\mathbf{x}}_{k}} = \begin{bmatrix} \left| \widetilde{x}_{k-1,1}(n) \right|^{2} & 0 \\ & \left| \widetilde{x}_{k-1,2}(n) \right|^{2} & \\ 0 & \ddots & \\ 0 & & \left| \widetilde{x}_{k-1,N}(n) \right|^{2} \end{bmatrix}$$

2)
$$\mathbf{W}_{k} = \left(\mathbf{SR}_{\tilde{\mathbf{x}}_{k}}\mathbf{S}^{H} + \hat{\mathbf{R}}_{v}\right)^{-1}\mathbf{SR}_{\tilde{\mathbf{x}}_{k}}$$

3)
$$\widetilde{\mathbf{x}}_{k}(n) = \mathbf{W}_{k}^{H}\mathbf{y}(n)$$

Termination:

 $\widetilde{\mathbf{x}}_k(n)$ vector converges.

3.3 SPECTRAL OVER-SAMPLING

As the name of the algorithm indicates, RISR is capable of achieving super-resolution under over-sampling. Super-resolution is defined as the ability to resolve two sources separated by less than the nominal resolution angle of $2\pi/N$ in electrical degrees under the nominal sampling of π/N . To incorporate spectral

over-sampling into the algorithm, the steering vector angle spectrum in the signal model can be over-sampled by a factor *P* and the angular spacing between the closest samples becomes:

$$\Delta \omega = \frac{\pi}{N} \cdot \frac{1}{P} \tag{3.8}$$

where *P* is the oversampling factor and P = 1 is equivalent to the nominal sampling. Signal vector **x** then becomes length 2P*N and **S** becomes a collection of 2P*N column steering vectors of length *N*.

The ability of the estimator to super-resolve signal sources is not a result of the over-sampling in signal angles. If an estimator cannot resolve angles at the nominal resolution, its performance remains the same even when the source angle space is over-sampled in the signal model.

The super-resolution probability as a function of the signal to noise ratio (SNR) with different values for over-sampling factor P are plotted for both uncorrelated and correlated signals in Chapter 5 under the RISR section. The results are also compared with the super-resolution performance of MUSIC.

3.4 MULTIPLE-TIME PROCESSING

The ability to estimate the DOA with only one snapshot signal sample is a major advantage that enables RISR to handle temporally correlated signals. Nevertheless, RISR can also be generalized to handle multiple snapshot data so that all the information contained in the collected data set can be extracted. Several multiple-snapshot processing schemes for increasing the SNR to boost the estimator performance were developed and are introduced in this chapter. To incorporate these schemes into the estimator, the underlying signals are assumed to be stationary within the time span of the received data.

3.4.1 INCOHERENT INTEGRATION

The first scheme, denoted as I2-RISR, incoherently integrates the estimates at the end of each stage in the algorithm. The main idea is to average the signal power estimates of each data snapshot at the end of the re-iteration stage before feeding it into the filter update for the next stage. Since the power of the estimates are averaged, phase information is lost and hence the term "incoherent" integration.

For the initialization stage and the first reiteration, each of the *K* snapshots (column vector) in the $N \times K$ matrix **Y** is processed independently the same way single-snapshot data is processed. Each column in **Y** is match-filtered with the steering vector **S** to create a vector of initial estimates. This is equivalent to the following matrix operation for the initialization stage:

$$\widetilde{\mathbf{X}}_0 = \mathbf{S}^H \mathbf{Y} \tag{3.9}$$

For each column in $\tilde{\mathbf{X}}_0$, denoted as a $\tilde{\mathbf{X}}_0^i$, a *first*-stage RMMSE filter W_1^i is created through the update of the structured covariance matrix $\mathbf{R}_{\tilde{\mathbf{X}}_0^i}$. The received data corresponding to that column is then filtered to form a vector of *first*-stage estimate $\tilde{\mathbf{X}}_1^i$. Before the *second* reiteration stage, the power of the *K* estimate $\tilde{\mathbf{X}}_1^i$'s are averaged in time to form the power estimates for the *first* stage:

$$\left|\overline{\mathbf{x}}_{1}(n)\right|^{2} = \left(\frac{1}{K}\right)\sum_{i=1}^{K}\left|\widetilde{\mathbf{X}}_{1}^{i}(n)\right|^{2}$$
(3.10)

The estimate $\bar{\mathbf{x}}_1(n)$ is then used to form the structured covariance matrix $\mathbf{R}_{\bar{\mathbf{x}}_2}$ needed to construct the *second*-stage I2-RISR filter:

$$\mathbf{W}_{2} = \left(\mathbf{S}\mathbf{R}_{\bar{\mathbf{x}}_{2}}\mathbf{S}^{H} + \mathbf{R}_{v}\right)^{-1} \left(\mathbf{S}\mathbf{R}_{\bar{\mathbf{x}}_{2}}\right)$$
(3.11)

where $\mathbf{R}_{\bar{\mathbf{x}}_{2}} = \begin{bmatrix} \left| \bar{x}_{1,1} \right|^{2} & 0 \\ \left| \bar{x}_{1,2} \right|^{2} & \\ & \ddots & \\ 0 & & \left| \bar{x}_{1,N} \right|^{2} \end{bmatrix}$

Notice only one filter is formed at the *second* stage. The multiple-snapshot data **Y** is then filtered by W_2 through matrix multiplication to form the *second*-stage estimates which undergoes a incoherent integration to form the vector of estimates $\bar{\mathbf{x}}_2(n)$ as the final output for the second reiteration:

$$\overline{\mathbf{x}}_{2}(n) = \left(\frac{1}{K}\right) \left|\widetilde{\mathbf{X}}_{2}\right| \cdot \left| \begin{array}{c} 1\\1\\\vdots\\1 \\ \end{bmatrix}_{N \times 1} \qquad \text{where } \widetilde{\mathbf{X}}_{2} = \mathbf{W}_{2}^{H} \mathbf{Y} \qquad (3.12)$$

The process repeats Equations (3.11) and (3.12) for subsequence stages until the solution $\overline{\mathbf{x}}_k(n)$ converges. Figure 3-2 demonstrates the block diagram of the overall operation:



Figure 3-2. Block diagram of I2-RISR operation

Note that the phase information of the estimate is lost due to the averaging of the power estimate at the end of each stage. However, it is not a concern since the goal is to determine the DOA and often times the magnitude of the signal component is sufficient for the signal estimation.

3.4.2 EIGEN DECOMPOSITION

The second scheme, denoted as e-RISR, has a much simpler formulation than the I2-RISR because it only differs from the original RISR by a pre-processing step which involves a transformation of the $N \times K$ multiple-snapshot data matrix into a $N \times 1$ data vector with single dimensional. This vector is then treated the same way as the observed input data vector $\mathbf{y}(n)$ in the original RISR for the rest of the processing. Assuming the data is stationary throughout the *K* snapshots, this scheme pre-combines the spatio-temporal data matrix **Y** into a spatial data vector by using the eigen decomposition to extract the spatial information common to all the data snapshots.

Let \mathbf{Y} be the matrix containing K snapshots of signal samples of length N. The dimension-reduced signal sample is obtained first by time-averaging the outer-products of each of the K snapshots:

$$\mathbf{C} = \left(\frac{1}{K}\right) \sum_{n=1}^{K} \mathbf{y}(n) \mathbf{y}^{H}(n) = \left(\frac{1}{K}\right) \mathbf{Y} \mathbf{Y}^{H}$$
where $\mathbf{Y} = \begin{bmatrix} \mathbf{y}(1) & \mathbf{y}(2) & \cdots & \mathbf{y}(K) \end{bmatrix}$
(3.13)

where the $N \times N$ matrix **C** is an approximation of the received signal covariance matrix. The **C** matrix is then eigen-decomposed to obtain a set of *N* length-*N* linearly independent eigenvectors **V** and a diagonal matrix **D** along which diagonal *N* corresponding eigen values are stored. They are then recombined to form a single data vector as shown in (17) below.

$$\widetilde{\mathbf{y}} = \sum_{i=1}^{N} \mathbf{V}_{i} \widetilde{d}_{i}$$
(3.14)
where $\mathbf{C} = \mathbf{V}^{H} \mathbf{D} \mathbf{V}, \ \widetilde{d}_{i} = \sqrt{\mathbf{D}_{ii}}$

The pre-processed vector $\tilde{\mathbf{y}}$ is a linear combination of the eigenvectors \mathbf{V}_i weighted by the corresponding square-rooted eigen values $\tilde{\mathbf{d}}_i$. This scheme is similar to time averaging the received signals except the outer products of snapshots are averaged to avoid signal cancellation due to phase differences over time. The vectors in $C^{\mathbb{N}}$ representing the dominating signal energy direction is enhanced because of the increase in SNR through the summation.

3.4.3 Additional Methods

A number of other approaches for incorporating multiple-snapshot data into RISR were also investigated; nonetheless the last scheme presented in this section is more meaningful. All methods were developed as an attempt to find the best representation of the original data matrix to maximize data extraction in the RISR algorithm. However, after comparing their performances from the Monte Carlo simulation results, which will be presented in Chapter 5, I2-RISR is chosen for RISR implementation throughout the rest of this thesis due to its consistency in performance under different underlying signal scenarios. I2-RISR is also simple to implement and analyze.

Integration with Auto-correlation Matrix as Filter Input

The operation of this approach differs from I2-RISR only in the input data matrix. Instead of the multiple-snapshot $N \times K$ received data matrix **Y**, the $N \times N$ autocorrelation matrix **C** is used for filtering. The initialization becomes:

$$\widetilde{\mathbf{X}}_{0} = \mathbf{S}^{H} \mathbf{C}$$
where $\mathbf{C} = \left(\frac{1}{K}\right) \mathbf{Y} \mathbf{Y}^{H} = \left(\frac{1}{K}\right) \sum_{n=1}^{K} \mathbf{y}(n) \mathbf{y}^{H}(n)$
(3.15)

The pre-averaged matrix at the k^{th} stage is then:

$$\widetilde{\mathbf{X}}_{k} = \mathbf{W}_{k}^{H} \mathbf{C}$$
(3.16)

To understand the matrix operation of this scheme more clearly, note that the i^{th} column of **C**, denoted as C_i , is the linear combination of the *K* data snapshots weighted by the i^{th} element of the corresponding snapshot, as shown in Equation (3.15). Therefore, the autocorrelation matrix **C** still contains all the spatio-temporal information in **Y**.

$$\widetilde{\mathbf{X}}_{k}^{i} = \mathbf{W}_{k}^{H} \mathbf{C}_{i}$$
where $\mathbf{C}_{i} = \left(\frac{1}{K}\right) \sum_{n=1}^{K} \mathbf{y}(n) \mathbf{y}_{i}^{*}(n)$
(3.17)

The rest of the algorithm for this scheme is identical to that of I2-RISR in Equations (3.11) and (3.12). Note that when the number of data snapshots K is greater than the size of antenna array, this scheme reduces the amount of the data RISR is fed to process.

CHAPTER 4: SOURCE AFFINE IMAGE RECONSTRUCTION

4.1 SAFFIRE ALGORITHM

4.1.1BASIC ALGORITHM

Source AFFine Image REconstruction (SAFFIRE) is based on the framework of the RMMSE approach developed in Section 3.1. The MEG dipole signal model presented in Chapter 2 is repeated here for convenience. Let *N* and *M* be the number of sensors and dipole grid, respectively. Equation (4.1) below shows the MEG dipole forward model:

$$\mathbf{y} = \mathbf{B} \, \mathbf{x} + \mathbf{v} \tag{4.1}$$

where **B** is the $N \times 2M$ leadfield matrix, **x** is the $2M \times 1$ dipole component strength vector, **v** is the $N \times 1$ additive noise vector and **y** is the $N \times 1$ vector of the MEG sensor signal corresponding to a single snapshot.

The derivation of the basic SAFFIRE algorithm is similar to that of RISR, which begins by substituting the MEG dipole model into the RMMSE filter formulation from Equation (4.2).

$$\mathbf{W}_{k} = \left(\mathbf{B}\mathbf{R}_{\widetilde{\mathbf{x}}(k)}\mathbf{B}^{H} + \mathbf{R}_{\mathbf{v}}\right)^{-1}\left(\mathbf{S}\mathbf{R}_{\widetilde{\mathbf{x}}(k)}\right)$$
(4.2)
where
$$\mathbf{R}_{\widetilde{\mathbf{x}}(k)} = \begin{bmatrix} \left|\widetilde{x}_{1}(k-1)\right|^{2} & 0\\ \left|\widetilde{x}_{2}(k-1)\right|^{2} & \\ 0 & \ddots\\ 0 & \left|\widetilde{x}_{2M}(k-1)\right|^{2} \end{bmatrix}$$
$$= \left(\widetilde{\mathbf{x}}(k-1)\widetilde{\mathbf{x}}^{H}(k-1)\right) \circ I_{2M \times 2M}$$

 $\mathbf{R}_{\tilde{\mathbf{x}}(k)}$ is the diagonal matrix of the signal power estimates from the previous iteration and $\tilde{x}_i(k)$ is the *i*th element of the estimated signal $\tilde{\mathbf{x}}(k)$. Zero values off

diagonal in $\mathbf{R}_{\tilde{\mathbf{x}}(k)}$ are from the assumption that signals from different directions are temporally uncorrelated. Therefore, the signal estimates at the k^{th} stage is computed as:

$$\widetilde{\mathbf{x}}(k) = \mathbf{W}_{k}^{H} \mathbf{y} = \mathbf{R}_{\widetilde{\mathbf{x}}(k)} \mathbf{B}^{H} \left(\mathbf{B} \mathbf{R}_{\widetilde{\mathbf{x}}(k)} \mathbf{B}^{H} + \mathbf{R}_{v} \right)^{-1} \mathbf{y}$$
(4.3)

Equations (4.2) and (4.3) serve as the core of the SAFFIRE algorithm with convergence generally attained within 10–15 iterations. However, because the forward model in Equation (4.1) is greatly underdetermined as discussed in Chapter 2 and the ill condition of the leadfield matrix **B** due to disparate attenuation effects resulting from different dipole depths relative to the brain surface, biasing effects would occur unless a proper initialization and solution scaling is used. To combat these biasing effects, the SAFFIRE algorithm operates in an affine transformed space and utilizes an initial estimate that is much less ambitious than the minimum-norm solution so as to avoid local minima. These specific characteristics of the SAFFIRE algorithm are described in the following sections.

4.1.2 AFFINE TRANSFORMATION OF SOLUTION SPACE

The ℓ^2 -norm of the individual leadfield vectors in **B** may vary considerably because the leadfield matrix incorporates the effects of attenuation induced by the source-to-sensors distance. As a result, minimum-norm initializations (such as used by FOCUSS) tend to produce initial estimates that are biased towards superficial dipole sources. To ameliorate this biasing problem the SAFFIRE algorithm utilizes an affine transformation of the solution space to remove the norm variations. The initial estimate of the dipole component strengths is then determined within this affine-transformed space.

The affine transformation is based on the equalization of the ℓ^2 -norms over the leadfield matrix. The transformation matrix **D** is formulated as

$$\mathbf{D} = \begin{bmatrix} \mathbf{b}_1^H \mathbf{b}_1 & & 0 \\ & \mathbf{b}_2^H \mathbf{b}_2 & & \\ & & \ddots & \\ 0 & & & \mathbf{b}_{2M}^H \mathbf{b}_{2M} \end{bmatrix}^{1/2} = \begin{bmatrix} |\mathbf{b}_1| & & 0 \\ & |\mathbf{b}_2| & & \\ & & \ddots & \\ 0 & & & |\mathbf{b}_{2M}| \end{bmatrix} = (\mathbf{B} \mathbf{B}^H) \circ \mathbf{I}_{2M \times 2M}$$
(4.4)

which is a diagonal matrix comprised of $|\mathbf{b}_i|$, the ℓ^2 -norms of the individual columns of the leadfield matrix **B**. It is mathematically equivalent to the Hadamard product of $(\mathbf{B}^{\mathrm{H}}\mathbf{B})^{1/2}$ and the $2M \times 2M$ identity matrix and hence it is invertible. Thus the affine transformation of the solution space is accomplished by re-expressing the forward model in Equation (4.1) as:

$$\mathbf{y} = \mathbf{B} \mathbf{x} + \mathbf{v}$$

$$= \mathbf{B} \mathbf{I} \mathbf{x} + \mathbf{v}$$

$$= \mathbf{B} \mathbf{D}^{-1} \mathbf{D} \mathbf{x} + \mathbf{v}$$

$$= \mathbf{B}_{\mathbf{a}} \mathbf{x}_{\mathbf{a}} + \mathbf{v}$$

where
$$\mathbf{B}_{\mathbf{a}} = \mathbf{B} \mathbf{D}^{-1} = \begin{bmatrix} \mathbf{b}_{1} & \mathbf{b}_{2} & \cdots & \mathbf{b}_{2M} \\ |\mathbf{b}_{1}| & |\mathbf{b}_{2}| & \cdots & |\mathbf{b}_{2M}| \cdot \mathbf{x}_{2M} \end{bmatrix}^{T}$$

and
$$\mathbf{x}_{\mathbf{a}} = \mathbf{D} \mathbf{x} = \begin{bmatrix} \mathbf{b}_{1} | \cdot x_{1} & |\mathbf{b}_{2}| \cdot x_{2} & \cdots & |\mathbf{b}_{2M}| \cdot x_{2M} \end{bmatrix}^{T}$$
(4.5)

The affine-transformed leadfield matrix B_a has unit column norms, and x_a contains the dipole component strengths scaled by the associated column norms of **B**.

Within the affine transformed space, the iterative estimation procedure of Equations (4.2) and (4.3) for the k^{th} stage can be applied by replacing **B** and $\tilde{\mathbf{x}}(k)$ with $\mathbf{B}_{\mathbf{a}}$ and $\tilde{\mathbf{x}}_{a}(k)$ in the formulation, respectively. After the terminal K^{th} iteration, the true estimate of the dipole component strengths in the original space can then be inversely transformed back from the transformed space by:

$$\widetilde{\mathbf{x}}(K) = \mathbf{D}^{-1} \, \widetilde{\mathbf{x}}_a(K) \tag{4.6}$$

4.1.3 MATCHED FILTER BANK INITIALIZATION

A good initial estimate is necessary for any recursive algorithms. For dipole localization problems, previous iterative approaches such as the FOCUSS algorithm

have employed the Moore-Penrose pseudo-inverse to obtain an initial estimate of the dipole component strengths as $\tilde{\mathbf{x}}_a(0) = \mathbf{B}_a^H (\mathbf{B}_a \mathbf{B}_a^H)^{-1} \mathbf{y}$, with the assumption of the affine-transformed space filter operation. However, due to severe ill-condition of the leadfield matrix **B**, the pseudo-inverse can bias the iterative solution to a basin of attraction for solutions closer to the MEG sensors as discussed in Chapter 2, thereby yielding estimates that are substantially different from the true solution. Once the recursive stage of the algorithm begins within the incorrect basin of attraction, the solution subsequently never converges to the true solution.

In contrast, the SAFFIRE algorithm utilizes a less ambitious approach to obtain the initial solution by employing a matched filter bank in the affine-transformed space to guarantee initial estimate in the correct basin of attraction:

$$\widetilde{\mathbf{x}}_{a}(0) = \mathbf{B}_{a}^{H} \mathbf{y}$$
(4.7)

Because the forward model is greatly underdetermined, the matched filter bank initialization provides rather poor spatial resolution due to the correlation between leadfield vectors that are in close spatial proximity.

The key to the effective SAFFIRE initialization method is that it utilizes 1) the affine-transformed \mathbf{B}_a instead of the filter containing ill-conditioned matrix $(\mathbf{B}_a \mathbf{B}_a^H)$ and 2) the matched filter bank to obtain an initial estimate. Without either of them, the initial solution would be biased towards incorrect basin of attraction. Note that a regularized version of the pseudo-inverse may also be used to initialize SAFFIRE as long as the regularization term λ is very large (often > 10⁴) to offset the ill-conditioning effects of $(\mathbf{B}_a \mathbf{B}_a^H)$:

$$\widetilde{\mathbf{x}}_{a}(0) = \mathbf{B}_{a}^{H} \left(\mathbf{B}_{a} \mathbf{B}_{a}^{H} + \lambda \mathbf{I} \right)^{-1} \mathbf{y}$$

$$\stackrel{\lambda \gg 10^{4}}{\approx} \mathbf{B}_{a}^{H} \left(\lambda \mathbf{I} \right)^{-1} \mathbf{y}$$

$$= \lambda^{-1} \mathbf{B}_{a}^{H} \mathbf{y}$$
(4.8)

Of course, as λ increases, the regularized pseudo-inverse converges to a scaled version of the matched filter. Thus, the preferred embodiment for SAFFIRE initialization is to employ the matched filter as in Equation (4.7).

The subsequent application of the iterative estimation process in Equations (4.2) and (4.3) of the SAFFIRE algorithm converges the solution to the individual active dipole components through concentrating the signal energy onto a particular subset of dipole components.

Despite of the low resolution of the initial estimate, the overall speed of convergence is only affected very slightly since from the second stage on, the algorithm is operated in affine-transformed space where the true solution can be quickly estimated by the SAFFIRE algorithm.

4.1.4 ENERGY NORMALIZATION

Recursive algorithms for any ill-conditioned system matrices requires special care for handling the energy or norm of the estimate signal vector at each stage due to the possibility of erroneous scale propagation and magnification through the recursive process, which, for SAFFIRE, could lead to zero value for all dipole strength estimates or even worse, filter operation instability.

Before any energy scaling was implemented, two problems were revealed during the analysis of the SAFFIRE algorithm because of the unreasonable dipole strength values. The first one was due to the matched filter bank estimate, which caused the norm of the initial estimate vector too large. This can be explained by the matched-filter-bank energy spread over a wide extent in the initial estimate solution space, as a result of the correlation between leadfield vectors in the sample space. The low resolution of the matched filter estimates visually shows this problem in Chapter 5.

The second energy-scaling problem originates from the recursive filter update formula in Equation (4.2). Substituting the affine-transformation space equation in (4.3) into the filter update equation we can obtain

$$\widetilde{\mathbf{x}}_{a}(k) = \mathbf{W}_{k}^{H} \mathbf{y} = \mathbf{R}_{\widetilde{\mathbf{x}}_{a}(k)} \mathbf{B}_{a}^{H} \left(\mathbf{B}_{a} \mathbf{R}_{\widetilde{\mathbf{x}}_{a}(k)} \mathbf{B}_{a}^{H} + \mathbf{R}_{v} \right)^{-1} \mathbf{y}$$
where $\mathbf{R}_{\widetilde{\mathbf{x}}_{a}(k)} = \begin{bmatrix} \left| \widetilde{\mathbf{x}}_{a,1}(k-1) \right|^{2} & \mathbf{0} \\ \left| \widetilde{\mathbf{x}}_{a,2}(k-1) \right|^{2} & \mathbf{0} \\ & \ddots & \\ \mathbf{0} & \left| \widetilde{\mathbf{x}}_{a,2M}(k-1) \right|^{2} \end{bmatrix}$

$$= \mathbf{D} \mathbf{R}_{\widetilde{\mathbf{x}}(k)} \mathbf{D}$$
(4.9)

where **D** is the affine-transform matrix and $\mathbf{R}_{\tilde{\mathbf{x}}(k)}$ is the Hadamard product of $\tilde{\mathbf{x}}(k) \tilde{\mathbf{x}}^{H}(k)$ and the identity matrix.

Manipulating the equation gives

$$\widetilde{\mathbf{x}}_{a}(k) = \mathbf{D} \, \mathbf{R}_{\widetilde{\mathbf{x}}(k)} \, \mathbf{D} \, \mathbf{D}^{-1} \, \mathbf{B}^{H} \left(\left(\mathbf{B} \, \mathbf{D}^{-1} \right) \mathbf{D} \, \mathbf{R}_{\widetilde{\mathbf{x}}(k)} \, \mathbf{D} \left(\mathbf{D}^{-1} \, \mathbf{B}^{H} \right) + \mathbf{R}_{\mathbf{v}} \right)^{-1} \mathbf{y}$$

$$= \mathbf{D} \, \mathbf{R}_{\widetilde{\mathbf{x}}(k)} \, \mathbf{B}^{H} \left(\mathbf{B} \, \mathbf{R}_{\widetilde{\mathbf{x}}(k)} \, \mathbf{B}^{H} + \mathbf{R}_{\mathbf{v}} \right)^{-1} \mathbf{y}$$

$$(4.10)$$

In order to understand the scaling problem, we first define the "estimated" sensor signal as the MEG data calculated using the forward model in Equation (4.1) by substituting in $\tilde{\mathbf{x}}_a(k)$ for the $\tilde{\mathbf{x}}(k)$

$$\widetilde{\mathbf{y}}(k) = \mathbf{B}_a \widetilde{\mathbf{x}}_a(k) \tag{4.11}$$

Substituting in Equation (4.10) yields

$$\widetilde{\mathbf{y}}(k) = \mathbf{B} \, \mathbf{D}^{-1} \, \mathbf{D} \, \mathbf{R}_{\widetilde{\mathbf{x}}(k)} \, \mathbf{B}^{H} \left(\mathbf{B} \, \mathbf{R}_{\widetilde{\mathbf{x}}(k)} \, \mathbf{B}^{H} + \mathbf{R}_{\mathbf{v}} \right)^{-1} \mathbf{y}$$

$$= \mathbf{B} \, \mathbf{R}_{\widetilde{\mathbf{x}}(k)} \, \mathbf{B}^{H} \left(\mathbf{B} \, \mathbf{R}_{\widetilde{\mathbf{x}}(k)} \, \mathbf{B}^{H} + \mathbf{R}_{\mathbf{v}} \right)^{-1} \mathbf{y}$$
(4.12)

The equation above indicates that a good dipole estimate $\tilde{\mathbf{x}}_{a}(k)$ should lead to very similar energy levels or norms between $\tilde{\mathbf{y}}(k)$ and \mathbf{y} . Observing Equation (4.12), if the column norms of $\mathbf{BR}_{\tilde{\mathbf{x}}(k)}\mathbf{B}^{H}$ are larger enough to dominate the inverse term, the product of the terms to the left of \mathbf{y} would become an identity matrix so that $\tilde{\mathbf{y}}(k)$ would approach \mathbf{y} . However, for reasons that will be discussed later, the norms of $\mathbf{BR}_{\tilde{\mathbf{x}}(k)}\mathbf{B}^{H}$ at the first few stages are a lot smaller than that of $\mathbf{R}_{\mathbf{v}}$, hence pre-multiplying \mathbf{y} by a matrix with small norms results in $\tilde{\mathbf{y}}(k)$ with smaller norm than y. The recursive nature of the algorithm drives the estimate $\tilde{\mathbf{x}}_{a}(k)$ to zero because of the diminishing energy level of the forward model estimate $\tilde{\mathbf{y}}(k)$ through the iterations.

To alleviate the energy scaling problems, SAFFIRE utilizes energy normalization at the end of each iteration. The energy normalization ensures that the dipole component estimate at a given iteration, when employed in the forward model (exclusive of noise), would yield a received signal estimate $\tilde{\mathbf{y}}(k)$ that possesses the same energy as the actual received signal \mathbf{y} . As a result, with an estimate of the received signal given $\tilde{\mathbf{x}}_a(k)$, the current estimate of the underlying dipole component strengths defined in Equation (4.9), the resulting energy estimate is determined as

$$\widetilde{\boldsymbol{\xi}}(k) = \widetilde{\mathbf{y}}^{H}(k)\widetilde{\mathbf{y}}(k)$$
(4.13)

Given the energy of the measured received signal as

$$\boldsymbol{\xi}_{\mathbf{y}} = \mathbf{y}^{H} \mathbf{y} \tag{4.14}$$

The energy-normalized dipole component strength estimate is thus

$$\widetilde{\mathbf{x}}_{a,norm}(k) = \sqrt{\frac{\xi_{\mathbf{y}}}{\widetilde{\xi}(k)}} \widetilde{\mathbf{x}}_{a}(k)$$
(4.15)

The energy normalization procedure as presented in Equations (4.13-15) is employed to the dipole component strength estimate at the end of each iteration of SAFFIRE.

Note that since leadfield vectors in **B** are highly correlated, the energy level of the matched filter or the low-resolution response is normalized to distribute the energy produced by a few true active dipoles among all the dipoles whose vectors are correlated with the true dipoles. Therefore, the elements in the dipole strength estimates are smaller than the actual underlying dipole strength. This is the reason that the energy level of $\mathbf{BR}_{\tilde{\mathbf{x}}(k)}\mathbf{B}^{H}$ is a lot less than that of $\mathbf{R}_{\mathbf{v}}$ at the matched-filter initialization as well as the first few stages.

4.1.5 NOISE COVARIANCE ESTIMATION

Unlike the FOCUSS algorithm, which requires the determination of a proper regularization term in the matrix inverse to accommodate the presence of additive noise in the forward model, SAFFIRE naturally incorporates that function into its algorithm through the signal model subsumed in the RMMSE formulation. The noise covariance matrix $\mathbf{R}_{\mathbf{v}} = E\{\mathbf{v} \ \mathbf{v}^H\}$ can be estimated directly as in Equation (4.16) below from the measured data over the interval of N_{noise} time samples in which no induced response is present. In so doing, the background noise as well as the ambient electromagnetic activity is captured, allowing the desired stimulated neural response to be isolated from the background activity.

$$\hat{\mathbf{R}}_{\mathbf{v}} = \frac{1}{N_{noise}} \sum_{k=1}^{N_{noise}} \mathbf{v}(n) \, \mathbf{v}^{H}(n)$$
(4.16)

4.1.6 IMPLEMENTATION

To summarize the SAFFIRE algorithm as discussed in previous subsections, the following outlines its principal components.

Implementation of the basic SAFFIRE algorithm

Initialization:

i) Compute estimate of noise covariance matrix over the interval of N_{noise} time samples that contains purely noise:

$$\hat{\mathbf{R}}_{\mathbf{v}} = \frac{1}{N_{noise}} \sum_{k=1}^{N_{noise}} \mathbf{v}(n) \, \mathbf{v}^{H}(n)$$

ii) Compute the measured received signal energy $\xi_{\mathbf{y}} = \mathbf{y}^{H}\mathbf{y}$ for the received signal \mathbf{y} at the time sample of interest.

iii) Compute the matched filter bank estimate of the affine-transformed dipole component strengths as

$$\widetilde{\mathbf{x}}_{a}(0) = \mathbf{B}_{a}^{H}\mathbf{y}$$
 where $\mathbf{B}_{a} = \mathbf{B}\mathbf{D}^{-1}$

iv) Energy-normalize the matched filter bank estimate as

$$\tilde{\mathbf{y}}(0) = \mathbf{B}_{a} \, \tilde{\mathbf{x}}_{a}(0)$$
$$\tilde{\xi}(0) = \tilde{\mathbf{y}}^{H}(0) \, \tilde{\mathbf{y}}(0)$$
$$\tilde{\mathbf{x}}_{a,norm}(0) = \sqrt{\frac{\xi_{y}}{\tilde{\xi}(0)}} \, \tilde{\mathbf{x}}_{a}(0)$$

Recursive stages:

For k = 1, 2, ..., K iterations, compute:

1) $\mathbf{R}_{\widetilde{\mathbf{x}}_{a}(k)} = \left(\widetilde{\mathbf{x}}_{a,norm}(k-1)\,\widetilde{\mathbf{x}}_{a,norm}^{H}(k-1)\right) \circ \mathbf{I}_{2M \times 2M}$

2)
$$\mathbf{W}_{k} = \left(\mathbf{BR}_{\widetilde{\mathbf{x}}_{a}(k)}\mathbf{B}^{H} + \hat{\mathbf{R}}_{\mathbf{v}}\right)^{-1}\mathbf{BR}_{\widetilde{\mathbf{x}}_{a}(k)}$$

- 3) $\tilde{\mathbf{x}}_{a}(k) = \mathbf{W}_{k}^{H}\mathbf{y}$
- 4) $\tilde{\mathbf{y}}(k) = \mathbf{B}_a \, \tilde{\mathbf{x}}_a(k)$

5)
$$\tilde{\boldsymbol{\xi}}(k) = \tilde{\mathbf{y}}^{H}(k) \tilde{\mathbf{y}}(k)$$

6)
$$\widetilde{\mathbf{x}}_{a,norm}(k) = \sqrt{\frac{\xi_{\mathbf{y}}}{\widetilde{\xi}(k)}} \, \widetilde{\mathbf{x}}_{a}(k)$$

Termination:

As $\tilde{\mathbf{x}}_{a,norm}(k)$ vector converges at the K^{th} stage, compute dipole component strength estimate as

$$\widetilde{\mathbf{x}} = \mathbf{D}^{-1} \widetilde{\mathbf{x}}_a(K)$$

4.1.7 RECONSTRUCTION OF DIPOLE TIME COURSE

Not only is SAFFIRE capable of estimating the location of activated dipole, the filter **W** as a result of the algorithm can also be used to reconstruct the underlying time-course of the estimated dipole for the sensor data outside of the processing time frame of the input data **y**, labeled as $\mathbf{y}(t)$, t = 1, ..., T, as shown in Equation (4.17).

The equation is applied with the assumption that $\mathbf{y}(t)$ consists of neural activity of only the estimated dipole since the reconstructed time course does not account for interference activities from other dipoles.

$$x_{j}(t) = \frac{\mathbf{w}_{K,j}^{H}}{|\mathbf{b}_{j}|} \mathbf{y}(t) \qquad \text{for } t = 1, ..., T$$

$$(4.17)$$

where *j* is the index of the estimated dipole, $\mathbf{w}_{K,j}$ is the *j*th column in the SAFFIRE filter from the last recursive stage, *K*, of the algorithm, and $x_j(t)$ is the estimated dipole strength at time index *t*. For the case where $\mathbf{y}(t)$ contains neural activity due to dipoles other than the estimated dipole, the reconstructed time course can be obtained by applying a modified filter, which is generated by SAFFIRE using data containing dipole activities not necessarily corresponding to consecutive time index. This approach will be discussed in greater details in the upcoming section.

4.2 MULTIPLE-TIME PROCESSING

Despite of the high temporal resolution SAFFIRE can achieve with only single sensor data snapshot, SAFFIRE can be generalized to process multiple data snapshots to maximize SNR so that more information can be extracted from provided data and potentially improve dipole localization accuracy. Another purpose of multiple-time processing is to reconstruct an interferer-free time-course for a particular dipole location. The multiple-time processing ability of SAFFIRE presented in this section is developed through the use of incoherent integration on estimated dipole strengths at each iteration.

For the purpose of increasing SNR, the underlying signals are assumed to be stationary within the time span of the received sensor data in order to enhance the underlying dipole component strength. However, the temporal resolution of the SAFFIRE algorithm is worsen as the time over which the data snapshot spans increases. Compared to other algorithms that require hundreds of sensor data snapshot to construct meaningful data covariance matrices, SAFFIRE can precisely localize an active dipole with typically around 5 data snapshots and as a result, its temporal resolution is still a great advantage over other dipole localization approaches.

The second way to make use of the multiple-time processing feature of SAFFIRE is to operate on sensor data snapshots corresponding to an underlying signal as well as on the snapshots containing other dipoles that are active at any time within the time-course of a dipole of interest. As such, the nature of the underlying signal responsible for the data snapshots is not restricted in terms of stationarity. Of course, the more stationary snapshots are processed, the better the ability of SAFFIRE is to localize the responsible active dipole. An example for the case involving interferer will be discussed at the end of the next chapter.

Let **Y** be a $N \times L$ matrix containing a collection of $N \times 1$ received signal (column) vectors over an interval of *L* time samples denoted as

$$\mathbf{Y} = \begin{bmatrix} \mathbf{y}(1) & \mathbf{y}(2) & \cdots & \mathbf{y}(L) \end{bmatrix}$$
(4.18)

Note that if the L time samples are consecutive, the incoherent integration in SAFFIRE algorithm results in SNR gain; while for the opposite case, SAFFIRE can be used as a tool for dipole time-course reconstruction. This latter implementation is useful for generating accurate EEG/MEG time courses that may consist of different spatial responses at different times.

The incoherent integration procedure is incorporated into the SAFFIRE algorithm as outlined at the end of the previous section in a similar fashion as I2-RISR. Steps ii), iii), and iv) of the SAFFIRE initialization are performed individually on each of the *L* time samples to obtain *L* initial matched filter bank estimates. These energy-normalized, affine-transformed dipole strength estimates are denoted as $\tilde{\mathbf{x}}_{a,norm,1}(0)$, $\tilde{\mathbf{x}}_{a,norm,2}(0)$, ..., $\tilde{\mathbf{x}}_{a,norm,L}(0)$ for the L time samples. The individual estimates are then combined incoherently in Step 1 of the first iteration (*k* = 1) as

$$\mathbf{R}_{\widetilde{\mathbf{x}}_{a,norm,1}} = \frac{1}{L} \sum_{i=1}^{L} \left\{ \left(\widetilde{\mathbf{x}}_{a,norm,i}(0) \, \widetilde{\mathbf{x}}_{a,norm,i}^{H}(0) \right) \circ \mathbf{I}_{2M \times 2M} \right\}$$
(4.19)

It is subsequently used to determine the MMSE-based filter bank \mathbf{W}_k in Step 2. This filter bank is then applied to each of the *L* received signal vectors $\mathbf{y}(1)$, $\mathbf{y}(2)$, ..., $\mathbf{y}(L)$ as in Step 3 followed by the *L* individual energy normalizations in Steps 4, 5, and 6. In general, the L estimate vectors obtained in Step 6 of a given iteration are combined in Step 1 of the following iteration as

$$\mathbf{R}_{\widetilde{\mathbf{x}}_{a,norm,\mathbf{k}}} = \frac{1}{L} \sum_{i=1}^{L} \left\{ \left(\widetilde{\mathbf{x}}_{a,norm,i}(k-1) \, \widetilde{\mathbf{x}}_{a,norm,i}^{H}(k-1) \right) \circ \mathbf{I}_{2M \times 2M} \right\}$$
(4.20)

4.3 MULTIPLE-STAGES FOR VOLUMETRIC CONSTRAINTS

The SAFFIRE algorithm has been found to perform very well at accurately identifying the spatial locations of brain activity regions. This spatial accuracy may be exploited to facilitate even greater accuracy by using the processed results to determine the volumetric region(s) of the brain in which activity is present during the time interval of interest. In so doing, the size of the leadfield matrix may be significantly reduced so as to only encompass a constrained region around the previous dipole estimate locations.

This matrix-size reduction is accomplished by replacing the leadfield matrix \mathbf{B}_{a} in the SAFFIRE algorithm, which comprises the entire sample region, with a modified leadfield matrix $\mathbf{B}_{a,r}$ that contains the subset of leadfield vectors corresponding to the constrained spatial region(s). Subsequently re-processing the received data with the volume-constrained SAFFIRE algorithm allows for much less spatial ambiguity such that further image accuracy can be achieved. Alternatively, the active spatial regions identified by SAFFIRE may be utilized as a priori knowledge for other EEG/MEG approaches that require such information [15, 40].

CHAPTER 5: SIMULATION RESULTS AND DISCUSSION

5.1 RISR

5.1.1 PERFORMANCE METRICS

The performance of RISR is demonstrated and analyzed for a number of simulation scenarios and is compared with MUSIC, SSMUSIC and FOCUSS in this section. The main metric used to evaluate algorithm performance is denoted as the probability of separation (POS), which is defined as follow: for the outcome of each trial of the Monte Carlo simulations, a value of "1" is assigned if (1) two individual peaks are identified within half the nominal resolution in both directions from the true null, which is defined as the mid-distance between the true sources; and (2) a 3 dB-null exists between the 2 estimated peaks; a value of "0" is assigned if any of (1) and (2) is not satisfied. For consistency, the angular spectrum is over-sampled by a factor such that 3 samples are between the two separate sources.

Although RISR is capable of estimating the source signal strength, the focus of application of this algorithm is to determine the direction from which signal arrives, and hence the mean square error (MSE) of the estimated signal strength, a common metric for estimator performance, does not serve as a metric in the scope of our discussion. This metric could also be somewhat misleading depending on what criteria the MSE calculation is based on.

However, a metric denoted as angular root mean square error (ARMSE) is defined to measure the amount of error by which an estimated angle is deviated from the true location. This is analogous to the MSE metric commonly used in DOA literature except it is predicated upon successful separation of two sources due to the fact that a single estimated peak does not correspond to any of the two true sources. The ARMSE in degrees is calculated as the square root of the average of the angular difference between the estimated and the associated true signal sources for the simulation trials assigned with a POS of 1.

5.1.2 BASIC PERFORMANCE

The performance of RISR algorithm with different processing schemes along with that of MUSIC and SS MUSIC algorithms are illustrated and compared in this section. Plots of RISR angular spectrum evolving over iterations, anecdotal results of a simple case setup, and several Monte Carlo simulation results are presented. The simulation setup of all cases consists of an uniform linear array (ULA) containing N = 10 antenna elements, each with half-wavelength spacing from adjacent elements, impringed by far-field signals. The nominal resolution and nominal sampling are hence $2\pi/N = 36^{\circ}$ and $\pi/N = 18^{\circ}$, respectively. The angular sample space is over-sampled by a factor of 4 to determine the probability of separation as well as for better visualization of resulting plots. Signals generated in all simulations are complex exponentials with unit magnitude and uncorrelated random phase uniformly distributed over 2π radians.

It is assumed that MUSIC and SS-MUSIC have the exact prior knowledge of the number of signal sources N_{source} and that SS-MUSIC divides the ULA into 5 subarrays with size of $N_{sub} = 6$ so that it satisfies the $N_{sub} \ge 2*N_{source}$ requirement. RISR is programmed to run K = 20 iterations after the matched-filter initialization to ensure convergence.

For the plots shown in Figure 5-1, single snapshot (K = 1) is taken for incoherent sources at SNR = 35 dB from angular electrical angles of -90°, -72° and 0°. The plot titled as "0" on the top left corner is the low-resolution matched filter response with peak energy levels around 15 dB. As the RISR iteration progresses through the first 8 RISR re-iterative stages until the spectrum converges to the sparse solution, which are illustrated in plots titled from "1" to "8" in Figure 5-1, the spectral energy level at peak angles increases while the attenuation of estimates from other angles deepens to the noise floor. Note that as the angular resolution increases from plot titled as "2" to "3", the sources with smaller angular distance are separated into two individual lobes with higher resolution. The trend in which the RISR angular spectrum converges to the true solution is similar for most cases of RISR





Figure 5-1. Angular spectrum (in electrical degrees) of RISR with K = 1 snapshot, N = 10, SNR = 35 dB and uncorrelated true sources at -90°, -72° and 0° over the first 8 iterations, including the initial solution titled as "0" on the top left corner

For the second anecdotal simulations, the basic performance of I2-RISR, e-RISR, RISR with covariance matrix **C** as input, MUSIC and SS-MUSIC are studied and compared through simple DOA simulation which consists of incoherent signals at SNR = 20 dB from electrical angles of -90°, -72° and 0°. A total of K = 20data snapshots were taken for the algorithms. The power spectrum of the signal estimates in dB are shown in Figure 5-2. Since the MUSIC and SS-MUSIC algorithms do not estimate underlying signal strength inherently, the scales used in the associated plots only reflect the presence of signal at the corresponding spectral location.



Figure 5-2. Angular spectrums of different algorithms with uncorrelated sources at -90° , -72° and 0° , K = 20 samples, $SNR = 20 \, dB$

As the angular spectra in Figure 5-2 illustrate, while all estimators are able to identify the all three sources, the three schemes of RISR achieve higher sparseness in the solution with substantially deeper nulls between the two closely-spaced sources than MUSIC and SS-MUSIC. There is more energy spread over the neighbor angles in the angular spectrum of SS-MUSIC than MUSIC because the use of sub-arrays for spatially smoothed covariance reduces the resolution of SS-MUSIC. For the case of e-RISR and I2-RISR with the covariance matrix **C** as data input, the estimated source angles are slightly deviated from the truth by an acceptable amount. However, they are not included in the final RISR algorithm because of the consistent performance of I2-RISR under the most circumstances. Since the original RISR is only a special case of I2-RISR with *K* = 1, RISR from here on refers to I2-RISR for simplicity.

Notice that the estimated values of the signal strengths for all three RISR estimators are very close to the actual signal strength of 1 (0 dB), demonstrating the accuracy of signal magnitude estimation the RISR can achieve. As for MUSIC and

SS-MUSIC, the spectral value in the spectrum has no indication of absolute or relative signal strength between signals from different angles.

Figure 5-3 to 5-6 illustrate the performance of the different methods with regard to the probability of separation of the nearby sources and the associated angular root mean square error. A total of K = 20 data snapshots are taken from a signal that consists of three uncorrelated sources at -90°, -81° and 0° and with 500 Monte Carlo runs are performed for each of the SNR value from 0 dB to 45 dB at 3 dB increments. The possible variations in the receive angle and possibility of the inter-sample source angles are accounted by independently assigning a random angle deviation uniformly distributed on $\Delta \theta \in \{-1^\circ, 1^\circ\}$ for each of the three sources for each Monte Carlo run. For clearer comparison, I2-RISR is compared to other multiple snapshot processing methods for RISR in Figure 5-3 and 5-4 while its results are displayed with MUSIC and SS-MUSIC in Figure 5-5 and 5-6.



Figure 5-3. Probability of separation of RISR, e-RISR and RISR with C as input versus SNR for uncorrelated sources at -90°, -81° and 0°, each with a random angle deviation $\Delta \theta \in \{-1^\circ, 1^\circ\}$, K = 20 samples

The superior performance of I2-RISR over other multiple snapshot processing schemes is demonstrated in Figure 5-3 above for uncorrelated sources. At an SNR of 21 dB, I2-RISR achieves a POS of 1 while the POS for e-RISR and RISR with

covariance matrix **C** as input are significantly lower at around 0.65 and 0.02, respectively. In order for e-RISR and the RISR with **C** methods to reach a POS of 1, they require SNR of 33 dB and 39 dB, respectively, which are a lot greater than the 21 dB for I2-RISR.



Figure 5-4. Angular root mean square error of RISR, e-RISR and RISR with C as input versus SNR for uncorrelated sources at -90°, -81° and 0°, each with a random angle deviation $\Delta \theta \in \{-1^\circ, 1^\circ\}$, K = 20 samples

ARMSE is defined only when nearby sources are separated and hence the ARMSE of the three RISR schemes as a function of SNR does not start until their corresponding POS values become non-zero. At the SNR value of 18 dB, the ARMSE of e-RISR starts off with a value roughly 2.3° smaller than I2-RISR. However, at the same SNR value in Figure 5-4, the POS value of I2-RISR is around 0.1 or 10% higher than e-RISR. At SNR value of 24 dB, the ARMSE of I2-RISR is at approximately 0.8°, significantly lower than that of the other 2 algorithms, which are 1.7° and 2.3°. Compared to the other schemes, I2-RISR consistently achieves lower ARMSE and higher POS over most SNR range.



Figure 5-5. Probability of separation of RISR, MUSIC and SS-MUSIC versus SNR for uncorrelated sources at -90°, -81° and 0°, each with a random angle deviation $\Delta \theta \in \{-1^\circ, 1^\circ\}$, K = 20 samples

Figure 5-5 shows that for a signal with SNR of 21 dB, RISR can successfully separate two close-by sources with a probability of 1 while MUSIC and SS-MUSIC algorithms have nearly zero probability in the same regard. At a probability of separation of 0.8, the SS-MUSIC and MUSIC algorithms require approximately 7 dB and 12 dB higher SNR than RISR, respectively. The reason for the inferior performance of MUSIC as compared to SS-MUSIC is that with the same amount of temporal snapshots, SS-MUSIC sub-divides them into overlapping snapshots for spatial-smoothing to exploit more information than original MUSIC does.


Figure 5-6. Angular root mean square error of RISR, MUSIC and SS-MUSIC versus SNR for uncorrelated sources at -90°, -81° and 0°, each with a random angle deviation $\Delta \theta \in \{-1^\circ, 1^\circ\}$, K = 20 samples

The non-zero value of ARMSE for SS-MUSIC at SNR = 15 dB in Figure 5-6 is only caused by the corresponding POS having a value slightly above zero as shown in Figure 5-5, and hence this is not taken into account when comparing the algorithm performances. The ARMSE of I2-RISR starts off at the SNR of 18 dB with a value around 0.6° smaller than that of SS-MUSIC and stays at a value smaller or equal to both MUSIC and SS-MUSIC algorithms over the entire SNR range. At SNR = 24 dB, ARMSE of I2-RISR is at around 0.5° while the other algorithms are at 1.1° and 1.2°. Compared to the other schemes, I2-RISR consistently achieves lower ARMSE and higher POS than MUSIC and SS-MUSIC algorithms over the SNR range in the simulation. In conclusion, from the discussion drawn from Figures 5-3 to 5-6, I2-RISR clearly out-performs all four other algorithms for uncorrelated signals.

5.1.3 TEMPORAL ROBUSTNESS

To demonstrate the signal correlation tolerance of RISR and the two algorithms under comparison, an anecdotal result and a Monte Carlo simulation results will be shown in this subsection. The anecdotal case carries the same setup as before except that the closely spaced signals from -90° and -81° are completely

correlated temporally. Signal correlation is modeled by restricting a phase difference between signals, where the phase difference is fixed for all K = 20 snapshots.



Figure 5-7. Anecdotal results of RISR, MUSIC and SSMUSIC with sources at -90°, -72° (correlated with source at -90°) and 0°, K = 20 samples, SNR = 35 dB

As shown in Figure 5-7, temporal correlation of underlying signals does not affect the performance of RISR. While RISR correctly estimates both the directions and the signal, MUSIC completely fails to detect the two coherent signals and provide a direction of estimated signal source midway between the two close-by underlying signals. SS-MUSIC detects the two coherent signals at the price of reduced angular resolution at the peaks in its angular spectrum because as explained in previous section, spatial-smoothing trades spatial resolution for temporal robustness through the use of overlapping sub-arrays. Note again that SS-MUSIC does not provide any information about the signal strength.

The temporal robustness of RISR is investigated through 500 Monte Carlo simulation run as illustrated in Figure 5-8 and 5-9. The simulation setup consists of three signals from electrical angles of -90°, -81° and 0, where the two near-by signals

are correlated in the same way as the previous anecdotal simulation except that the phase difference between them is randomized for each Monte Carlo run. Individual random angle deviation $\Delta \theta \in \{-1^\circ, 1^\circ\}$ is also assigned to each of the three sources for each Monte Carlo run. A total number of K = 20 snapshots are taken. The SNR for the simulation varies from 18 to 60 dB with an increment of 3 dB.



Figure 5-8. Probability of separation of RISR, MUSIC and SS-MUSIC versus SNR for correlated sources at -90°, -81° and an uncorrelated at 0°, each with a random angle deviation $\Delta \theta \in \{-1^\circ, 1^\circ\}$, K = 20 samples

It can be observed from Figure 5-8 that MUSIC completely fails to separate signal sources as expected when correlation between signals is present. In order to achieve a POS of 0.8 or 80%, RISR requires a SNR value of approximately 30 dB, which is 13 dB lower than the 43 dB required for SS-MUSIC. At *SNR* = 30 dB, the POS of RISR reaches 75%, whereas that of SS-MUSIC is only 5%. Compared to the uncorrelated case shown in Figure 5-5, SS-MUSIC is affected more significantly by signal correlation than RISR. For instance, at POS of 80%, the SNR required for SS-MUSIC rises to 43 dB for correlated case from 28 dB for uncorrelated case while for RISR it only increases to 30 dB from 20 dB. The reason for the performance degradation for SS-MUSIC is that for uncorrelated case, spatial-smoothing increases the amount of data for constructing the data covariance matrix and thus it exploits more information about the signal sources; whereas for correlated signals, that

functionality of spatial-smoothing is replaced by making the data-covariance full-rank to break down the signal correlation.



Figure 5-9. Angular root mean square error of RISR, MUSIC and SS-MUSIC versus SNR for correlated sources at -90°, -81° and 0°, each with a random angle deviation $\Delta \theta \in \{-1^\circ, 1^\circ\}$, K = 20 samples

As shown in Figure 5-9, at SNR = 18 dB the ARMSE of SS-MUSIC and RISR are of 12° and 2.5°, respectively. As SNR increases, the ARMSE of RISR decreases very slightly and is at roughly 2° while that of SS-MUSIC converges to zero. The small non-zero ARMSE for RISR occurs only for a certain higher super-resolution factor values. This effect is a function of signal separation distance and is an estimate bias due to a certain range of phase difference between the two close-by sources, which causes signal cancellation in the matched filter estimate and biases the sparse source convergence in the RISR iterations.

Although SS-MUSIC performs slightly better than RISR in terms of ARMSE, at a lower SNR values as shown in Figure 5-8, RISR achieves significantly higher POS, which is the more meaningful metric in the scope of DOA estimation. In addition, RISR is not sensitive to the number of sources whereas, assuming the knowledge of number of sources P, SS-MUSIC limits the number of subarray elements to be more or equal to 2P. With an antenna array size of N = 10, SS-MUSIC breaks down if the number of signal sources surpasses five.

5.1.4 SPECTRAL SUPER-RESOLUTION

The nominal spectral resolution is conventionally defined as $2\pi/N$, where *N* is the number of sensor elements. The ability of RISR to super-resolve spectral sources is presented and compared with SS-MUSIC algorithm, which is known to have super-resolution capability. Monte Carlo simulation with 500 runs is performed for a setup with N = 10, K = 40 data snapshots, and two uncorrelated sources at -90° and $-90^\circ + \Delta\theta_L$, where $\Delta\theta_L = \frac{2\pi}{N \cdot L}$, for a range of super-resolution factor $L = \{2, 4, 8, 6, 8, 16\}$. The two source angles are assigned an independent random angle deviation uniformly distributed on $\Delta\theta \in \{-1^\circ, 1^\circ\}$ at each Monte Carlo run to account for angle deviation for the received signal. The results for POS and ARMSE are shown in Figure 5-10 and 5-11, respectively, for the two algorithms.



Figure 5-10. Monte Carlo simulation result of POS for two uncorrelated signals over a range of super-resolution factor L versus SNR for RISR and SS-MUSIC algorithms

Observe in Figure 5-10 that when the sources separation is close to the nominal resolution with lower *L* values, RISR and SS-MUSIC have similar POS across all SNR. As the super-resolution factor *L* increases, which decreases the separation $\Delta\theta_L$ between the source angles, the performance of SS-MUSIC degrades more significantly than RISR. More specifically, when *L* goes from a factor of 2 to 4, the SNR required for RISR to achieve a POS of 80% increases from 14 dB to 20 dB,

while that for SS-MUSIC increases from 14 dB to 25 dB. Observe from the figure that for L = 8 (black curves), the POS of RISR approaches 90% at around SNR = 26 dB, but for the same POS value SS-MUSIC requires 12 dB higher SNR than RISR. For the extreme case of L = 16 where the separation between the sources is only 2.25°, the POS of RISR and SS-MUSIC are 100% and 0%, respectively, at SNR value of 33 dB. This Monte Carlo simulation shows that RISR out-performs SS-MUSIC in super-resolution in all simulated cases except for low SNR and small super-resolution factors L, in which case SS-MUSIC requires a few dB less than RISR to achieve a low POS.



Figure 5-11. Monte Carlo simulation result of ARMSE for two uncorrelated signals over a range of super-resolution factor L versus SNR for RISR and SS-MUSIC algorithms

Signal sources start to be resolved at various SNR for the different super-resolution factors *L*, hence the ARMSE curves do not begin at the same SNR value as shown in Figure 5-11. The overall pattern of the two plots suggests that for correlated sources estimates of RISR have less angular error than that of SS-MUSIC at lower SNR and similar angular error at higher SNR. For example for L = 4, the ARMSE of RISR at 18 dB SNR is 2.5°, almost 3° smaller than SS-MUSIC and the ARMSE values are very close at 42 dB for both algorithms. Similarly, at SNR = 30 dB, the ARMSE of RISR is roughly 1.2° lower than that of SS-MUSIC for L = 16 while at SNR = 42 dB, the ARMSE values are both about 0.4°.

5.1.5 DATA SAMPLE SUPPORT

The goal of the next Monte Carlo simulation is to investigate how the amount of received data samples affects the performance of RISR. Similar to earlier sections, the simulation setup consists of three uncorrelated source from angular electrical angles of -90°, -81° and 0° with individual random angle deviation $\Delta \theta \in \{-1^\circ, 1^\circ\}$ assigned to each source for each Monte Carlo run. For each value of *K*, the number of data snapshots, 500 Monte Carlo runs are performed. The six *K* values simulated are 1, 2, 4, 8, 16 and 500. Note that RISR with *K* = 1 is equivalent to the original RISR with single data snapshot. The simulation results of RISR and SS-MUSIC are with regard to their probability of separation are shown in Figures 5-12 and 5-13.



Figure 5-12. Probability of separation of RISR versus SNR for uncorrelated sources at -90°, -81° and 0° and K = 1, 2, 4, 8, 16 and 500 snapshots



Figure 5-13. Probability of separation of SS-MUSIC versus SNR for uncorrelated sources at -90°, -81° and 0° and K = 1, 2, 4, 8, 16 and 500 snapshots

With a large amount of data support where the number of data snapshots K = 500, SS-MUSIC out-performs RISR as indicated by the corresponding POS curves (in cyan) in Figure 5-12 and 5-13. SS-MUSIC reaches a POS of 80% at SNR = 17 dB but RISR requires 2 dB higher SNR to achieve the same POS. As the number of snapshots K is reduced to 16, the ability to separate sources for SS-MUSIC degrades significantly while the POS of RISR is only minimally affected. The POS of SS-MUSIC at SNR = 20 dB decreases from 95% to 5% for a sample support reduction from K = 500 to 16 snapshots, whereas the POS of RISR remains at 90% for both K values. The intolerance of low sample support for SS-MUSIC can be observed by further comparing the POS curves. In order to achieve a POS of 60%, SS-MUSIC requires SNR values of 26 dB, 28 dB, 33 dB and 40 dB for K = 16, 8, 4 and 2 snapshots, respectively, which are a lot higher than the corresponding SNR values for RISR which are 18 dB, 18 dB, 18 dB and 20 dB, respectively. The insensitivity of the algorithm performance to sample support size K is an advantage of RISR over SS-MUSIC.

For the extreme case with only one single data snapshot (K = 1), the POS of SS-MUSIC are roughly zero even at high SNR, indicating SS-MUSIC fails to

separate signal sources under very low data sample support. RISR, on the other hand, achieves over 90% of POS at SNR = 35 dB. The ability of estimating signal sources correctly with only one data snapshot is one of the major advantages of RISR over SS-MUSIC and can be very useful when estimation is performed in a fast changing environment where only small data support is available [41]. Note that despite RISR is inferior to SS-MUSIC for the case of large data support, the stationarity required by SS-MUSIC limits the amount of data collected in DOA estimation, and hence K = 500 snapshots is rarely possible in most scenario.

5.1.6 CALIBRATION ERROR

An unavoidable error in real-world DOA applications created by physical antenna configuration is the calibration error, which refers to the discrepancy between the antenna characteristics accounted in the measurements and the actual values. The effects of calibration error on RISR and SS-MUSIC are investigated in this sub-section through Monte Carlo simulations for four cases: calibration error-free, 5% amplitude error only, 5% phase error only and both amplitude and phase errors. The signal consists of three uncorrelated sources from -90°, -81° and 0° and K = 40 snapshots are sampled. For each of the SNR values, 500 Monte Carlo runs are performed to obtain the plots in Figure 5-14 below.



Figure 5-14. Effect of calibration error on RISR and SS-MUSIC at each antenna element for three uncorrelated sources at -90°, -81° and 0°, N = 10 and K = 40 snapshots for amplitude error of 5% and phase error of 5%

Comparing the plots for the two algorithms in Figure 5-14, the degradation of SS-MUSIC due to phase error (black and red curves) at antenna elements is more severe than that of RISR. Observe that the POS of SS-MUSIC never exceeds 0.2 when phase error is present, whereas RISR achieves 80% of POS at SNR = 20 dB for both cases with phase error. As indicated by the green curves, SS-MUSIC also suffers more than RISR for signals with amplitude error. For example, at SNR = 21 dB, POS of RISR reaches 100%, about 70% more than the POS of SS-MUSIC. The inability of SS-MUSIC to separate nearby sources can be explained by the eigen method that MUSIC is based on. As a result of the small variation of the signal at each antenna element, the dominant eigenvectors of the original error-free data covariance matrix is represented as the combination of two eigenvectors, one with higher energy level than the other, in the new data covariance matrix. Although information of the underlying signal is contained in both eigenvectors, MUSIC includes the eigenvector with less energy or eigenvalue in the noise space, through which part of the signal is projected out from the solution. Therefore, SS-MUSIC breaks down when calibration error is present.

Note that for the cases containing phase errors at antenna sensors, the degradation of RISR at SNR above 33 dB is due to line-splitting. The two signals at -90° and -81° are estimated as three peaks in close angular proximity in the angular spectrum. The reason for this phenomenon is that at lower SNR, RISR considers the small variation/error on antenna elements as noise, whereas at higher SNR, the phase error of the antenna elements is more "noticeable" and thus causes the line-splitting of RISR to account for the error.

The amount by which POS is influenced by calibration error at antenna elements for each algorithm can be observed from the Monte Carlo simulation results shown in Figure 5-15 below. The simulation setup involves K = 40 snapshots sampled from a signal containing three uncorrelated sources from -90°, -81° and 0° at SNR = 25 dB. For each of the amplitude error varying from 0% to 10% of the unit-magnitude signal, 500 Monte Carlo runs are performed for each of the 1%

interval. Monte Carlo simulation is also performed for phase error in the same range of error percentage values where the phase error is uniformly distributed within the percentage of the 2π interval centered around 0 radian.



Figure 5-15. Probability of separation versus percentage calibration error in amplitude (left) and in phase (right) for RISR and SS-MUSIC with three uncorrelated sources at SNR = 25 dB and K = 40 snapshots

Observe from Figure 5-15 that as the amplitude and phase calibration errors percentage increase, the probability of separation of SS-MUSIC worsens. The POS value of SS-MUSIC drops from 0.8 for the error-free case to 0.5 for a 6% amplitude error and to almost 0 for the same amount of phase error, which indicates that the performance of SS-MUSIC is more sensitive to phase error than amplitude error. SS-MUSIC algorithm starts to break down in the presence of even very small phase error. On the other hand, the POS of RISR stays around 1 for the range of error under test in both plots as shown in Figure 5-15, suggesting the robustness of RISR to both types of calibration error.

5.1.7 COLORED ADDITIVE NOISE

The additive noise buried in received signal is not always angularly uniform in all directions and hence its noise spectrum is not completely "white" [42, 43]. To demonstrate the effect of colored additive noise to RISR and two other algorithms, an anecdotal simulation result of this scenario is presented in Figure 5-16. The received signal at SNR = 15 dB consists of three uncorrelated signals from -90°, -72° and 0°.

The colored noise was generated by filtering a spatially "white" signal through a bandpass filter with the mainlobe centered at 45°. A number of K = 8 data snapshots are taken for processing and $K_{noise} = 50$ snapshots of the pure-noise signal are used to obtain the noise covariance matrix $\mathbf{R}_{\mathbf{v}}$.



Figure 5-16. Angular spectrum of estimated signal strengths for different algorithms with 3 uncorrelated signals from -90°, -72° and 0° and addictive noise with mainlobe at around 45°

As shown in the angular spectrum at the bottom left plot in Figure 5-16, SS-MUSIC estimates only the signal source at 0° correctly. As mentioned before, SS-MUSIC assumes a white noise spectrum as well as the knowledge of the number of signal sources, which is three in this simulation, and thus it merges the two sources at -90° and -72° together as one estimated signal in order to account for the dominant energy at 45° in the signal space of the covariance matrix. Although the formulation of FOCUSS resembles that of RISR, the Least Squares approach does not inherently incorporate the presence of noise into the system model. Without taking noise into account, the output of the FOCUSS estimator is corrupted by an estimate of the

source signal at a noise-concentrated angular direction. Since RISR assumes the presence of noise in its formulation, the knowledge of the additive noise can be built into the algorithm. As a result, RISR is able to detect the correct signal sources in the presence of colored noise with a higher success rate than SS-MUSIC and FOCUSS.

5.1.8 OBSERVATION OF SPARSE SOLUTION CONVERGENCE

The recursive, non-linear nature of RISR makes developing the convergence proof for the algorithm a challenging task, despite of the successful identification of signal sources by RISR in a large number of scenarios presented in the Monte Carlo simulation results in previous sections. However, sparse solution convergence of RISR can be demonstrated by observing the dominant energy spread in the eigen spectrum of the filter after the solution of RISR has converged. A few anecdotal cases are shown in this section to exemplify the convergence of the number of dominant eigenvalues to the number of signal sources. Since the filter from Equation (3.4) is not a square matrix, the signal model equation (3.1) is substituted into the formula for the k^{th} estimate to obtain

$$\widetilde{\mathbf{x}}_{k} = \mathbf{R}_{\widetilde{\mathbf{x}}_{k}} \mathbf{S}^{H} \left(\mathbf{S} \mathbf{R}_{\widetilde{\mathbf{x}}_{k}} \mathbf{S}^{H} + \hat{\mathbf{R}}_{\mathbf{v}} \right)^{-1} \left(\mathbf{S} \mathbf{x} + \mathbf{v} \right)$$

$$\approx \mathbf{R}_{\widetilde{\mathbf{x}}_{k}} \mathbf{S}^{H} \left(\mathbf{S} \mathbf{R}_{\widetilde{\mathbf{x}}_{k}} \mathbf{S}^{H} + \hat{\mathbf{R}}_{\mathbf{v}} \right)^{-1} \mathbf{S} \mathbf{x} = \mathbf{P} \mathbf{x}$$
(5.1)
where $\mathbf{P} = \mathbf{W}^{H} \mathbf{S}$

P is the $M \times M$ modified filter matrix of which the eigen spectrum will be observed in this section. Observe that as k gets large, the eigen space of matrix **P** is expected to converge to the a sparse representation.

The first anecdotal case involves a single signal at -90° with SNR = 20 dB and K = 20 snapshots of received data with the angular spectrum over-sampled by a factor of 10. The setup of the next two cases are identical to the first except two uncorrelated signals at -90° and -45° and three uncorrelated signals at -90°, -45° and 0° are present for the second and third cases, respectively. Figure 5-17 below shows the 10 most dominant eigen values of **P** at the 15th iteration for all three cases.



Figure 5-17. Ten most dominant eigen values of matrix **P** at the 15th iteration for the estimation of (a) a single signal source at -90°; (b) two uncorrelated signals at -90° and -45°; (c) three uncorrelated signals at -90°, -45° and 0° with $SNR = 20 \ dB \ and \ K = 20$

Observe from Figure 5-17 that the number of dominant eigen values in the eigen spectrum of \mathbf{P} for each of the three cases equals the corresponding number of underlying signal sources. This observation demonstrates that the eigen spectrum of the solution space of RISR algorithm converges to the most sparse representation, which is a function of the number of distinct signal sources. Though a proof is not available, the convergence for RISR is strongly suggested by the three anecdotal examples.

5.2 SAFFIRE

5.2.1EXPERIMENTAL SETUP AND MEG CONFIGURATIONS

The performance of SAFFIRE is evaluated using computer simulations, which allow an objective assessment of the dipole localization accuracy for a number of dipole activation scenarios. Several examples of dipole source reconstruction are illustrated in the figures in this section to demonstrate the advantages of SAFFIRE compared to other schemes in terms of their performance. The simulation setup of dipole localization experiments is based on the physical setup of the MEG sensor equipments at the Hoglund Brain Imaging Center at the University of Kansas Medical Center. The human brain volume is evenly divided into 9014 sample grids for dipole localization. The MEG signals received at the 150 sensors of the

biomagnetometer system are simulated for different dipole activation scenarios at a sampling rate of 600 Hz. Figure 5-18 below illustrates the MEG sensor geometrical configuration with respect to a brain volume.



Figure 5-18. Physical configuration of 150 MEG sensors relative to the brain. Sensors are axial gradiometers with 2 detector coils separated by 5 cm. For better visualization, only the lower coil of each sensor is shown.

For each of the 9014 grids throughout the sample space, the leadfield vectors for only the phi and theta components (in spherical coordinates) of the associated unit dipole are computed due to the fact that the magnetic fields generated by the radial component are undetectable outside of the brain in the spherically-symmetric approximation of the volume conductor [21]. Hence, the size of leadfield matrix **B** responsible for dipole activity simulation is 150-by-18028. Each dipole is labeled a value from 1 to 9014 corresponding to the specific grid in the sample space.

SAFFIRE algorithm used for all the simulation results incorporates second-stage volumetric constrained processing after the initial SAFFIRE algorithm

is performed. The volumetric constrain to the sample space is implemented by first selecting a group of "active" dipole with at least 30% of the maximum estimated strength from the original SAFFIRE, then the final set of dipoles is formed by keeping all dipoles within 2 cm proximity of each dipole from the group of active dipoles. The leadfield vectors corresponding to the dipoles in the final set is concatenated to construct \mathbf{B}_{ar} , the volumetrically constrained leadfield matrix.

5.2.2 SINGLE DIPOLE ACTIVATION

In this section, the performance of SAFFIRE for single dipole activations will be investigated for three dipoles of various depths in the sample space of the brain. For all three simulation cases, a dipole activation curve of 2-second duration peaks at t = 1 s of the activation with a peak dipole strength of 30 nAm and a pulse width of 150 ms. The activation curve shown in Figure 5-19 below is sampled at 600 Hz which means a total of 1200 samples are collected with a peak at the 600th sample.



Figure 5-19. Activation curve for single dipole activation

The corresponding MEG sensor signal in matrix form of size 150×1200 , **Y**, is generated using the forward model, which involves multiplying a dipole activation row vector **x** by the leadfield vector of the theta component of the activated dipole \mathbf{r}_i , $\mathbf{b}(\mathbf{r}_i, \theta)$, where the product is then added to a noise vector **v**. The mathematical operation is presented in Equation (5.2).

$$\mathbf{Y}_{150 \times 1200} = \mathbf{b}(\mathbf{r}_i, \theta) \,\mathbf{x} + \mathbf{v} \tag{5.2}$$

where the sizes of **x**, $\mathbf{b}(\mathbf{r}_i, \theta)$ and **v** vectors are 1×1200 , 150×1 and 150×1 , respectively. The additive noise magnitude is 10 femto-Tesla which, for example, results in a SNR of the MEG signal at the peak MGFP to be around 17.44 dB for the superficial dipole in the first single dipole simulation case. This SNR belongs to the lower end of the typical SNR range of neural signals.

An example of the MEG sensor measurements at the 150 channels associated with single dipole activation is displayed in the lower plot of Figure 5-20 and the averaged power of the signals across sensors, also known as mean global field power (MGFP) is included in the upper plot of the same figure. The received data sample with maximum MGFP power is used as the input data for dipole localization of SAFFIRE algorithm. The criterion for SAFFIRE to estimate the active dipole is by selecting dipoles with 20% or more of the maximum estimated dipole power throughout the source space. It is observed that volumetric constrained reprocessing of SAFFIRE only improves the estimates for single dipole activation slightly.



Figure 5-20. Mean Global Field Power (MGFP, at the top) and the sensors response of the simulated MEG measurements (at the bottom) corresponding to single dipole activation.

Single Dipole Activation, eccentricity of 6.1 cm

The first simulation is performed for a superficial dipole labeled as 8981, which is 6.1 cm from the center of the brain. Figure 5-21 shows the true dipole in red as well as the estimated active dipole (in blue) by different algorithms in the 3D brain map. The input data matrix **Y** is simulated using Equation (5.2) with the leadfield vector corresponding to the theta component of dipole 8981. The first algorithm of which the performance is evaluated is SAFFIRE with single snapshot (K = 1) at the 600th sample of the simulated data. SAFFIRE is repeated to obtain the next set of results for multiple-time processing with K = 4 snapshots, which is associated with data from t = 995 ms to t = 1 s. Since FOCUSS and MNE algorithms were only developed for single snapshot data, only the 600th sample is used for the third and fourth set of results.







(b)



Figure 5-21. The angled (right) and front/top (left) view of the location of the true active dipole 8981 (in red), which is 6.1 cm from the center of the brain, estimated (in blue) by (a) SAFFIRE using K = 1 snapshots at 600th sample; (b) SAFFIRE using K = 4 snapshots centered around 600th sample; (c) FOCUSS using the 600th sample; and (d) MNE (in blue) using the 600th sample with a threshold of 70% of maximum estimated dipole strength

Comparing the results from (a) and (b) in Figure 5-21, both algorithms accurately estimate the location of the active dipole, indicating that SAFFIRE with single snapshot data is able to localize the superficial dipole as well as SAFFIRE with multiple-snapshot processing. FOCUSS in (c) also successfully estimate the superficial dipole correctly. However, determining the parameters for regularization

and norm un-biasing in the FOCUSS algorithm in Equation (2.29) require considerable amount of effort because they are functions of a number of factors such as the active dipole location and noise level. Due to the low-resolution nature of MNE algorithm, a threshold of 70% of maximum estimated dipole strength still results in a 3D dipole estimation map with very poor spatial resolution. MNE is clearly inferior to other algorithms in terms of spatial precision of dipole localization and is hence not considered in further performance comparison for algorithms. MNE is also expected to fail for inner dipoles due to the lack of any scheme for norm-bias adjustment.



Figure 5-22. Dipole strength estimated by SAFFIRE (top left), the neural activity index resulted from LCMV algorithm (top right), and the dipole strength estimated by MNE (bottom) for activated dipole 8981

The plots in Figure 5-22 represent the relative dipole strengths estimated by different algorithms. Observe that MNE algorithm delivers a "spectrum" with the worst resolution compared to the other two algorithms as expected. LCMV requires 400 snapshots, an equivalent of 0.667 s duration of data, to construct the data covariance matrix while only four snapshots, which translates to only 0.0067 s, are processed for the SAFFIRE algorithm, suggesting the higher temporal resolution capability of SAFFIRE in dipole localization. A norm-bias adjustment of each dipole for LCMV is achieved through the neural activity index (NAI), which is the ratio between the biased estimate of the variance (energy) of the dipole and the estimate of variance of the noise generated by the same dipole. As shown in Figure 5-22, the plot of dipole strength estimates by SAFFIRE achieves a remarkably higher level of sparseness than the NAI plot of LCMV, thus demonstrating the superior spatial resolution of SAFFIRE. In addition, the greater algorithm efficiency of SAFFIRE can be easily concluded from the fact that the computation time of LCMV was about 4 minutes, which is 20 times of the 12 seconds required by SAFFIRE.



Figure 5-23. True activation curve (in green) of dipole 8981 and reconstructed time course of dipole activity (in blue) by (a) SAFFIRE algorithm with K = 4 snapshots and (b) LCMV with 400 snapshots to construct the data covariance matrix

The reconstructed time course of dipole 8981 for SAFFIRE with K = 4 snapshots and for LCMV algorithms are plotted in blue in Figure 5-23. Observe that compared to LCMV, the time course for SAFFIRE matches slightly better with the

true activation curve plotted in green and maintains a lower noise level throughout the entire time span, suggesting that SAFFIRE can reconstruct cleaner time courses.

Single Dipole Activation, eccentricity of 4.4 cm

The second simulation is performed for a mid-depth dipole labeled as 7952, with a distance of 4.4 cm from the center of the brain. Figure 5-24 shows the true dipole in red as well as the estimated active dipole by different algorithms in the 3D brain map. Again, the MEG sensor data is simulated using Equation (5.2) with the corresponding data for dipole 7952. The algorithms of which results are shown in Figure 5-24 are SAFFIRE with single snapshot (K = 1), SAFFIRE with multiple-time processing (K = 4), and FOCUSS with single snapshot data and same algorithm parameters as the previous simulation.







Figure 5-24. The angled (right) and front/top (left) view of the location of the true active dipole 7952 (in red), which is 4.4 cm from the center of the brain, estimated (in blue) by (a) SAFFIRE using K = 1 snapshots at 600th sample; (b) SAFFIRE using K = 4 snapshots centered around 600th sample; and (c) FOCUSS using the 600th sample

The depth of dipole 7952 creates greater attenuation of magnetic fields, which consequently worsens the SNR in MEG sensor signal **Y** than that of dipole 8981 in the previous simulation. The 3D plot shown in (a) and (b) of Figure 5-24 illustrates that the SNR gain due to multiple data snapshots enables SAFFIRE to accurately localize the mid-depth dipole when single-snapshot SAFFIRE can only estimates the rough location of the same dipole. On the other hand, FOCUSS completely fails to localize the active dipole and instead, only provide active dipole estimates at superficial locations as shown in (c). This can be explained by the lack of robust bias adjustment of the dipole norm and the need to "tune" the parameters of the FOCUSS algorithm to their optimal values depending on the characteristics of the underlying dipole. These factors limit the performance of FOCUSS whereas SAFFIRE is free of such constraints for estimation involving non-superficial dipoles because of the affine-domain operation and energy normalization at each iteration.



Figure 5-25. True activation curve (in green) of dipole 7952 and reconstructed time course of dipole activity (in blue) by (a) SAFFIRE algorithm with K = 4 snapshots and (b) LCMV with 400 snapshots to construct the data covariance matrix

The reconstructed time course by SAFFIRE shown in Figure 5-25(a) for dipole 7952 is more noisy than that shown in Figure 5-24(a) for dipole 8981. The same can also be observed for the time courses by LCMV in Figure 5-24(b) and Figure 5-25(b). The reason for that is, as mentioned previously, the attenuation of magnetic field as it travels through the brain increases with dipole depth which causes the MEG signal SNR to decrease. Compared to LCMV, SAFFIRE produces a reconstructed time course with a lower noise level over the entire duration as well as a better-matched pulse around t = 1s.

Single Dipole Activation, eccentricity of 2.0 cm

The last single active dipole simulation is performed for a deep dipole labeled as 3310, which is 2.0 cm from the center of the brain. Figure 5-26 shows the true dipole in red as well as the estimated active dipole by different algorithms in the 3D brain map. Again, the MEG sensor data **Y** of size 150×1200 is simulated using Equation (5.2) with the corresponding data for dipole 3310. The algorithms of which results are shown in Figure 5-26 are SAFFIRE with single snapshot (*K* = 1) and SAFFIRE with multiple-time processing (*K* = 4).



Figure 5-26. The angled (right) and front/top (left) view of the location of the true active dipole 3310 (in red), which is 2.0 cm from the center of the brain, estimated (in blue) by (a) SAFFIRE using K = 1 snapshots at 600th sample; and (b) SAFFIRE using K = 4 snapshots centered around 600th sample

SAFFIRE with K = 4 snapshot is able to localize accurately an active dipole very deep in the brain as shown in Figure 5-26(b) whereas with only a single data vector, SAFFIRE can still estimate the approximate location of dipole 3310. Although SAFFIRE requires four snapshots to achieve high performance, the unprecedented temporal resolution of this algorithm demonstrated by accurate estimation using only an equivalence of 6.67 ms duration of data is a great advantage over other existing algorithms.



Figure 5-27. Dipole strength estimated by SAFFIRE on the left and the neural activity index resulted from LCMV algorithm on the right for activated dipole 3310



Figure 5-28. True activation curve (in green) of dipole 3310 and reconstructed time course of dipole activity (in blue) by (a) SAFFIRE algorithm with K = 4 snapshots and (b) LCMV with 400 snapshots to construct the data covariance matrix

Performance degradation of LCMV is illustrated by comparing the NAI plot of LCMV for dipole 3310 in Figure 5-27 to that for dipole 8981 in Figure 5-22. The energy is spread out among wider range of dipoles for the deeper dipole, which effectively lower the spatial resolution of LCMV. SAFFIRE, however, maintains the sparseness in the solution and return a dominant energy estimate at the correct dipole location as shown in Figure 5-27, indicating that SNR of sensors signal only affects the performance of SAFFIRE minimally for single dipole activation. As expected, the reconstructed time courses of dipole 3310 shown in Figure 5-28 has higher noise level than the corresponding curves for dipole 7952 in Figure 5-25 for the respective algorithms due to SNR deterioration. Observe that SAFFIRE reconstructed a time course with significantly lower the noise level and a more defined pulse curve than the LCMV algorithm. From the comparisons drawn for the three single dipole activation simulations, SAFFIRE undoubtedly out-performs LCMV in dipole time course reconstruction.

5.2.3 NEARBY DIPOLE PAIR

In this section, the performance of SAFFIRE is demonstrated for a dipole simulation of the theta components of a pair of superficial dipoles, which are 1.09 cm apart and are labeled as 8570 and 8629. Both dipoles have identical activation curve to the single dipole simulation case and MEG data matrix is simulated using Equation (5.2) with the data corresponding to the dipole pair. SAFFIRE is performed with K = 4 snapshots centered around the 603rd sample and volumetric constrained reprocessing. The sensor data is also processed by LCMV to yield a NAI plot for comparison with SAFFIRE.



Figure 5-29. Dipole strength estimated by SAFFIRE on the left and the neural activity index resulted from LCMV algorithm on the right for the dipole pair 8570 and 8629



Figure 5-30. Angled (left) and front (right) 3D-view of dipoles 8570 and 8629 (in red), which are 1.09 cm apart and about 5 cm from the center of the brain, and dipole estimated by SAFFIRE (in blue) using K = 4 snapshots centered around 603^{rd} sample



Figure 5-31. Reconstructed time course of dipole 8570 (left) and 8629 (right) using SAFFIRE (in blue) and the true dipole activation curve (in red)

It can be observed in Figure 5-29 that the NAI values of the majority of the dipoles in the NAI plot of LCMV is around 16.5 and the dipoles associated with the peak NAI do not match with the correct values of 8570 and 8629, indicating that LCMV fails completely to localize two close-by fully-correlated sources.

On the contrary, the plot of the dipole strength estimated by SAFFIRE shown in Figure 5-29 contains two peaks at the correct active dipole index of 8570 and 8629 while displaying the high spatial resolution by maintaining the sparseness in the solution "spectrum". Moreover, the spatial resolution of active dipoles achieved by SAFFIRE is demonstrated in Figure 5-30, which illustrates accurate localization of the two close-by dipoles. SAFFIRE also successfully reconstructs the time courses for the two active dipoles with small noise energy as illustrated in Figure 5-31. The ability to correctly separate close-by dipole with high spatial resolution as well as to reconstruct time course proves the distinctly advanced performance of SAFFIRE over LCMV.

5.2.4 MIRRORED DIPOLE-PAIR IN PRIMARY AUDITORY REGIONS

LCMV is known to break down when spatial correlation or interference is present in the signal and although the modified version, known as MCMV-CSRS, can solve such problem, it requires prior knowledge of the location of the interferer. In this section, the ability of SAFFIRE to handle spatial correlation without any prior knowledge of the underlying dipole activity is evaluated through the simulation of a pair of spatially-mirrored dipole located in the primary auditory regions. The dipoles are labeled as 1088 and 7952 and are both roughly 4.5 cm from the center of the brain. The activation curve of both dipoles are identical to the single dipole simulation case and MEG data matrix **Y** is simulated using Equation (5.2) with the data corresponding to the dipoles.



Figure 5-32. Angled (left) and front (right) 3D-view of the dipoles in the primary auditory regions (in red) and dipole estimated by SAFFIRE (in blue) using K = 4 snapshots centered around 600th sample



Figure 5-33. Reconstructed time course of dipole 1088 (left) and 7952 (right) in the auditory region using SAFFIRE (in blue) and the true dipole activation curve (in red)

The spatial correlation robustness of SAFFIRE algorithm is demonstrated through observations from Figure 5-32 and 5-33. As shown in Figure 5-32, the locations of both active dipoles are accurately estimated by SAFFIRE. The time course of the dipole in each primary auditory region reconstructed by SAFFIRE matches with the true underlying activation curve with small amount of noise as illustrated in Figure 5-33. The ability for SAFFIRE to handle the case with mirrored-dipoles in the primary auditory regions is the motivation for testing SAFFIRE for the ASSR simulation, of which the results are presented next.

5.2.5 AUDITORY RESPONSE

Auditory onset response is an evoked auditory response (EAR) in the primary and secondary cortex due to brief auditory stimuli presented at short intervals [15]. Studies suggest that EAR can be potentially used for hearing threshold or cognitive evaluation and measuring the consciousness level of patients undergoing anesthesia. Accurate reconstruction of the underlying spatio-temporal process of the primary and secondary cortex is essential to the studies of the neuromagnetic EAR and the effects induced by other factors. EAR involving short time delay of activation between dipole pairs in primary and secondary cortex requires very high level of spatio-temporal resolution and as a result, the reconstruction of this type of response had been a challenging problem in the neuroimaging community with conflicting results due to the different source reconstruction strategies employed by different studies.

The configuration of ASSR setup is shown in Figure 5-34 below with two dipole pairs, which are numbered as 1088, 7952, 613 and 8570, mirrored along the mid-sagital plane. One mirrored dipole pair is simulated in the regions of the primary auditory cortex and another in the secondary auditory cortex. The separations between dipoles 1088 and 613 in the left cerebral hemisphere and that between dipoles 7952 and 8570 in the right cerebral hemisphere are 1.52 cm and 1.65 cm, respectively. As illustrated in Figure 5-35, the dipole activation curves of all four dipoles are identical except that the peaks of activation curves for the secondary cortex are delayed by 50 ms, or 30 samples, from those in the primary cortex. Peak dipole strength is 30 nAm and the noise magnitude is 10^{-15} T.



Figure 5-34. Locations of the dipole pairs in primary cortex (red) and secondary cortex (green) for the ASSR simulation



Figure 5-35. Activation curves of primary and secondary auditory regions with peaks at t = 1 and 1.05 s (top left), the mean global field power of the received signals (top right) and the MEG sensors response of all 150 channels (bottom left)

Methods

The MEG sensor signals sampled at 600 Hz are generated by summing the MEG responses corresponding to the activation curve of each of the four dipoles computed using Equation (5.2). SAFFIRE algorithm is used in a manner similar to an overlapped sliding window for the data set of 78 samples from t = 962 ms (578th) sample) to t = 1.09 s (655th sample). At each estimation interval, K = 4 consecutive data snapshots are processed to yield an estimate and hence producing 75 estimates. Besides the original SAFFIRE algorithm, SAFFIRE with volumetric-constrained reprocessing as well as 3-of-5 detection are also carried out at each estimation interval for performance comparison. For simplicity, SAFFIRE with volumetric-constrained reprocessing is interchanged with SAFFIRE-2 in the rest of this section.

In order to remove the few occurrences of improbable estimates such as split dipoles or active dipole that only occurs for 1 sample (1.7 ms), the 3-of-5 detection is applied, also as an overlapped sliding window, to the output of SAFFIRE-2 and is defined as follows: for each group of five consecutive estimates, if a dipole considered as active (with at least 30% of the maximum estimated strength) for three or more of those five estimates, the dipole is counted as active. Since for each group of five estimates, one 3-of-5 detector estimate is produced, the total number of estimates after applying 3-of-5 detection is 71, four less than that of the other two methods.

Results

Three stages of estimated dipole activity are identified after observing the temporal progression of the dipole estimate locations for each of the three methods. The first and third stage corresponds to the period when the true dipole pair in the primary cortex and secondary cortex, respectively, is successfully localized. The second stage, which typically takes place between the peaks of the primary and secondary activation curves, is the transition interval where the dipoles estimated at locations between the primary and secondary cortices.

It is found from the activation curves that the period over which the power of the dipole associated with the primary activation is at least 20% of its maximum is from t = 968 ms (582nd sample) to t = 1.032 s (620th sample), whereas the period corresponding to the secondary activation is from t = 1.018 ms (612rd sample) to t = 1.082 s (650th sample). In order to compare the performance of the three different reconstruction methods, the ideal values of time indices will be referred to, which are listed in Table 5-1 below. Note that the ideal time period for the second stage is when the primary and secondary activation curves overlap.

	Ideal time	
First stage begins	968 ms (582)	
Second stage begins	1.018 s (612)	
Third stage begins	1.032 s (620)	
Third stage ends	1.082 s (650)	

Table 5-1. Ideal time indices for the first to third stages of ASSR estimation. The bracket value is the sample index corresponding to the time

The time index at which each stage occurs for the three methods are tabulated in Table 5-2. Since the estimates in the first or third stage might not be completely accurate or exact, a "soft" criterion is defined that the localization of a dipole is determined to be successful only when the estimated dipoles are exactly at or very close to the true dipole location with no ambiguity that it is in the correct location.

Table 5-2. Estimated time of ASSR dipole localization for the three processing schemes where the bracket value is the sample index corresponding to the time

	SAFFIRE	SAFFIRE-2	3-of-5 detector
First stage begins	980 ms (589)	968 ms (582)	968 ms (582)
Second stage begins	997 ms (599)	1.002 s (602)	1.002 s (602)
Third stage begins	1.045 s (628)	1.045 s (628)	1.043 s (627)
Third stage ends	1.077 s (647)	1.088 s (654)	1.080 s (652)

Observe from Table 5-2 that the time indices defining the boundary of the three stages matches better with the ideal case for SAFFIRE-2 than for the original SAFFIRE. The SAFFIRE-2 and ideal time index for the beginning of the first stage are both 582, which are 7 snapshots before the corresponding time index for SAFFIRE. Compared to the ideal duration of 8 snapshots for the second stage, SAFFIRE-2 with 26 snapshots is closer to ideal than SAFFIRE with 29 snapshots.

Although 3-of-5 detector provides very little improvement over SAFFIRE-2 in terms of the time indices for defining the stages, it delivers cleaner 3D dipole localization images during the second stage where the dipoles "migrate" from primary to secondary cortex and during the time period outside of any of the three stages due to low SNR. Three example sets of the dipole localization 3D plots corresponding to each of the three stages are demonstrated in Figures 5-36, 5-37 and 5-38 below along with the MGFP plots for results of which 3-of-5 detector is applied.



Figure 5-36. An example of the first stage. Top: true dipoles (in red) and the dipole locations estimated by SAFFIRE-2 with 3-of-5 detector (in blue) in 3D plot for data at around t = 972 ms (584th sample); bottom: MGFP plot from t =968 ms to t = 1.085 with primary and secondary activation curves which are out of scale. The black vertical line indicates t = 972 ms on the time axis



Figure 5-37. An example of the second stage. Top: true dipoles (in red) and the dipole locations estimated by SAFFIRE-2 with 3-of-5 detector (in blue) in 3D plot for data at around t = 1.025s (616th sample); bottom: MGFP plot from t = 968 ms to t = 1.085 with primary and secondary activation curves which are out of scale. The black vertical line indicates t = 1.025s s on the time axis


Figure 5-38. An example of the third stage. Top: true dipoles (in red) and the dipole locations estimated by SAFFIRE-2 with 3-of-5 detector (in blue) in 3D plot for data at around t = 1.052s (632^{rd} sample); bottom: MGFP plot from t =968 ms to t = 1.085 with primary and secondary activation curves which are out of scale. The black vertical line indicates t = 1.052 s on the time axis

Observe the MGFP plot within the estimation period in Figure 5-38 that the power received at MEG sensors at the peak of the primary activation is less than that at the peak of the secondary activation. This is due to more attenuation associated with the greater depth of the dipole pair in the primary cortex and hence lower SNR for the primary activation. Therefore, even though primary and secondary activation curves are identical with small delay, the third stage is estimated with a longer duration than the first stage as illustrated in Table 5-2. For the same reason, the duration of the first stage corresponding to the estimated primary activation is slightly shorter than that that of the third stage.

Figure 5-36 and 5-38 illustrate the accurate dipole localization of the dipole pairs in the primary and secondary cortices, respectively by SAFFIRE-2 with 3-of-5 detector. The transitional stage example shown in Figure 5-37 associated with data does not display correct localization of either dipole pairs, the estimates are still within the auditory cortex in the respective cerebral hemispheres. The temporal progression of the 3D plot during the transition stage shows the gradual migration of estimated active dipole from the primary cortex to secondary cortex.

In conclusion, the high temporal and spatial resolution of SAFFIRE enable the successful reconstruction of the spatio-temporal signal for the ASSR, which can motivate further neural signal processing to observe other types of neural dynamic scenarios.

Videos displaying the temporal progression of the dipole localization in 3D plots for ASSR as well as the three single dipole activation examples are created and can be requested through email.

5.2.6 TIME-COURSE RECONSTRUCTION WITH INTERFERENCE

Interference, such as energy from other dipole sources, maybe present within the temporal window of the reconstructed time course, yet outside of the time range over which data snapshots are taken for constructing the dipole localization filter. The ability of SAFFIRE to remove such interference with an additional multiple-time processing is demonstrated through a simulation of two mirrored dipoles of which the pulses of the dipole activation curves are separated with no overlapping.

The activation curves of the two dipoles in the primary auditory cortex, numbered as 1088 and 7952, and the MGFP of the MEG sensor signal are shown in Figure 5-39. The pulse of dipole 7952 is delayed by 500 ms, or 300 samples, from the pulse of dipole 1088 so that when K = 4 snapshots corresponding to the peak of MGFP at t = 1 s (601st sample) are processed by SAFFIRE, the activity of the interferer dipole (7952) does not contribute to dipole strengths estimation. The resulting 3D plot and the reconstructed time course of the estimated active dipole are shown in Figure 5-40.



Figure 5-39. Activation curves of dipoles 1088 and 7952 in the primary cortex with peak delay of 500 ms (left) and the MGFP plot of the MEG sensor signal (right)



Figure 5-40 Angled (top left) and front (top right) 3D-view of dipoles 1088 and 7952 in the primary cortex (red) with 500 ms peak delay and the dipole estimated by SAFFIRE (blue). The bottom plot is the reconstructed time course of the estimated dipole 1088 with a false peak at 1.5 s (blue) and the true activation curve (red)

Since the data snapshots around t = 1 s consisting of only the activity of dipole 1088 is selected to construct the filter, SAFFIRE identifies active source at dipole 1088 though not dipole 7952 as shown in Figure 5-40. The reconstructed time course of dipole 1088 contains a large peak around t = 1 s as expected and a smaller peak at t = 1.5 s. If no prior knowledge of the underlying dipole activity is available, the smaller peak could be a correct dipole pulse but it could also be a false peak due to activation of other dipoles which are spatially correlated with dipole 1088.



Figure 5-41. Angled (top left) and front (top right) 3D-view of dipoles 1088 and 7952 in the primary cortex (red) with 500 ms peak delay and the dipoles estimated by SAFFIRE (blue). The bottom plots are the reconstructed time course of the dipoles 1088 and 7952 (blue) and the true activation curve (red)

The multiple-time processing feature of SAFFIRE can be utilized in an alternative way to determine the true nature of the second peak. Four additional data snapshots around the time index associated with the second peak, i.e. t = 1.5 s (902nd sample), are concatenated with the original four snapshots around t = 1 s to form a new set of mixed input data for the SAFFIRE algorithm. The 3D plots of the estimated dipoles from the SAFFIRE reprocessing with mixed input are shown in Figure 5-41 along with the two reconstructed time courses.

With the MEG data input containing activities of the two dipoles, SAFFIRE accurately localized the dipoles responsible for the mixed sensor signal as shown in the upper plots of Figure 5-41. The time courses of the estimated dipoles in the bottom plots match very well with the associated underlying activation curves. Observe that the second peak of the reconstructed time course at t = 1.5 s in Figure 5-40 does not appear in the corresponding plot in Figure 5-41, whereas the reconstructed time course of the dipole 7952 has a peak at t = 1.5 s. The observations imply that the second peak in Figure 5-40 was indeed a false peak containing dipole energy of dipole 7952 due to the spatial correlated of the two dipoles. SAFFIRE filter can reconstruct dipole time courses only with information embedded in the input data the filter is constructed with. Intuitively speaking, since the mixed data carries information of the interferer, SAFFIRE is able to project out the dipole energy leakage of dipole 7952 from the reconstructed time course of dipole 1088 while the first SAFFIRE fails in the same regard. The results of this simulation case demonstrate the ability of SAFFIRE to successfully reconstruct spatially uncorrelated time courses through a secondary processing with input data containing activities of potential interferers.

CHAPTER 6: CONCLUSIONS AND FUTURE WORK

6.1 Conclusions

The novel algorithm developed for direction of arrival estimation, denoted as Re-Iterative Super-Resolution (RISR), recursively update the filter through the power estimates in the formulation until the solution converges to a maximally sparse solution. The results in Chapter 5 shows that RISR out-performs the algorithms under comparison in both spatial and temporal resolution. Super temporal resolution enabled by the single-snapshot processing becomes a major advantage in the presence of fast-moving targets, whereas finer spatial resolution increases the success of distinguishing closely-separated sources by the ULA. The temporal correlation robustness of RISR is a useful property when multiple signal sources are synchronized. Calibration error at each of the antenna elements is inevitable in antenna arrays setup and therefore the tolerance of RISR toward such error demonstrated through simulations is extremely useful in DOA estimation with real data. The susceptibility to colored noise or interference as well as the ability to estimate signal strength in addition to the aforementioned advantages reflects the great potential of the RISR algorithm.

Source AFFine Image REconstruction, an iterative approach based on the MMSE framework, is a powerful tool for functional neural imaging with magnetoencephalography signals as shown in the results of a number of dipole localization simulations. However, SAFFIRE can be applied to inverse modeling with other imaging techniques, such as magnetocardiography for the heart [44] and magnetogastrography [45] for the abdomen, provided that accurate forward models exist. Parametric neural dipole localization algorithms are known for heavy bias in favor of superficial dipoles. SAFFIRE was shown to produce bias-free estimates in simulations with activated dipoles of various depths without the need for parameter adjustment. The spatial resolution achieved by SAFFIRE due to solution sparseness confines the activate dipole region and, hence, is helpful for separating two close-by

dipoles. The exceptional temporal resolution is another advantage of SAFFIRE that allows the observation of neural evoked response dynamics in finer details. The unprecedented spatio-temporal resolution, temporal correlation robustness, independence of statistical prior knowledge and the effective dipole activity time course reconstruction of SAFFIRE together build the success in reconstructing the multiple sources of auditory evoked responses which is a difficult problem in the functional imaging community.

In spite of the similarities between the algorithms, such as the framework on which they are based and their recursive natures, RISR and SAFFIRE are two distinct approaches with a few principal differences. First of all, SAFFIRE is specific for inverse modeling of the biological process while RISR is applied to the more general DOA estimation. The signal model of RISR assumes far-field signals impringing on the ULA whereas the conformed array antenna configuration of SAFFIRE does not make such assumption. Lastly, SAFFIRE incorporates energy normalization and affine-transformed space operation at each iteration into the algorithm to alleviate the effects of matrix ill-conditioning, which is not a problem with the DOA estimation for the RISR algorithm.

6.2 Future Work

The evaluation of RISR algorithm shown in Chapter 5 was based on the simulation of sources with equal signal strengths. Therefore, other aspects of the algorithm, such as the signal sources dynamic range, distributed sources scenarios and real data processing, can be the next steps of investigation. The RISR is built upon a MMSE framework with a two-dimensional signal model for the azimuth angle. An extension of RISR into three-dimensional DOA estimation can be explored by considering a signal model that incorporates azimuth and elevation of the source signals.

The purpose of applying a 3-of-5 detector to a dipole activation response reconstructed by SAFFIRE is analogous to smoothing the spatial dipole estimates

across time in order to eliminate "short-lived" estimates, due to the low physical probability of such activation duration. This temporal averaging of spatial information is an example of space-time coupling processing, which enables information extraction from the relationship between the current and additional dimensions. Instead of post-processing, the first order information between space and time, such as the slope of activation curve of a detected dipole, might possibly be incorporated into the filter formulation to increase the accuracy of the overall reconstructed response.

SAFFIRE was shown to accurately localize dipole activities and reconstruct time course of the dipole activities by the results of a limited number of scenarios. Therefore, other scenarios, such as dipole activation curve shape and distributed dipole activity, can be simulated to examine the performance of the algorithm. Also, SAFFIRE is still yet to be tested on real experimental MEG data of the auditory evoked responses.

In spite of the disadvantages of other medical imaging techniques as discussed in the background section, some related research has been conducted for fMRI constrained MEG signal processing with other algorithms [46]. This approach should be investigated further since other imaging methods offer complementary information that will potentially increase the performance of SAFFIRE.

The sensor signals in the forward model from which SAFFIRE is derived can be substituted with ones related to other physical phenomena, for example, electroencephalography, magnetocardiography and magnetogastrography. The unprecedented spatio-temporal resolution of SAFFIRE algorithm can benefit the estimation of the underlying activities with higher accuracy and precision.

The applications of the recursive MMSE framework in radar adaptive pulse compression, direction of arrival estimation and functional neural imaging have been shown to achieve excellent performance. While the algorithms, depending on the application, might require different adjustment, the common formulation of the basic RMMSE filter should allow flexible application to other types of signal estimation.

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