

Trine Nyholm Kragh & Laura Nyrup Mogensen Mathematical Engineering, MATTEK

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STUDENT REPORT

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Bayesian Dictionary Learning for EEG Source Identification

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Participant(s):

Trine Nyholm Kragh Laura Nyrup Mogensen

Supervisor(s):

Jan Østergaard Rasmus Waagepetersen

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Abstract:

The thesis deals with the problem of recovering original brain source signals from low-density EEG measurements. Based on state of the art methods, an algorithm was proposed, for reproducing the current results. gorithm aim to leverage a covariancedomain dictionary learning (Cov-DL) method and a multiple sparse Bayesian learning (M-SBL) method. The proposed application of Cov-DL did not succeed. Thus, an alternative solution was proposed. The proposed algorithm was tested on EEG scalp measurements and compared to solutions obtained by ICA of high-density EEG measurements. Furthermore, a frequency analysis was performed comparing raw measurements and the recovered sources. From the perspective of practical use, an estimation of the unknown number of active source signals was proposed. It was concluded that the proposed algorithm did not reproduce the state of the art results. Though the M-SBL method alone was successful and a potential was seen for an estimation of the number of active sources, from M-SBL.

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Preface

Here is the preface. You should put your signatures at the end of the preface.				
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Danish Summary

I dette kandidatspeciale undersøges der hvordan man kan lokalisere og identificere den direkte hjerne aktivitet ud fra EEG-målinger. Dette omtales som at løse det inverse EEG-problem hvor det ønskes at finde en matrix bestående af hjerneaktiviteten - sources - samt en mixing matrix som tilføjer støj til sources. Det ønskes at løse det inverse EEG-problem for EEG-målinger hvor flere sources er tilstede end sensorer. Der vil tages udgangspunkt i to "state of the art" metoder, covariance-domain dictionary learning (Cov-DL) og multiple sparse Bayesian learning (M-SBL), til at løse dette problem. Ud fra disse to valgte metoder vil deres reproducerbarhed blive undersøgt for at se om samme konklusion opnås, at disse metoder effektivt løser det inverse problem. I dette projekt vil disse to metoder indgå i en samlet metode, Cov-DL vil bruges til at finde mixing matricen og M-SBL benytter sig af den fundne mixing matrix til at finde source matricen. Sekundært vil der undersøges i at modificere den samlede metode ud fra et praktisk perspektiv. Ved implementering af den samlede metode vil de enkle metoder blive testet og analyseret på simuleret data for at undersøge om metoden lykkes. Til dette vil de fundne estimater blive sammenlignet med de rigtige estimater ved brug af mean-square-error (MSE). For Cov-DL lykkes det ikke at genskabe den samme konklusion som den videnskabelige artikel bag metoden så denne har ikke en tilstrækkelig grad af reproducerbarhed. Den anden metode, M-SBL, findes vellykket når den sande mixing matrix benyttes. Herfra kan det konkluderes, at den tilsvarende videnskabelige artikel tilvejebragte en tilstrækkelig grad af reproducerbarhed. For at kunne anvende den samlede metode på rigtig EEG-måling så vælges der at erstattet Cov-DL med en fast mixing matrix som er fundet ud fra empiriske tests. Med denne ændring og ud fra tests af den samlede metode så forventes det ikke at der opnås en tilstrækkelig identificering og lokalisering af source matricen. Da de sande estimater ikke kendes for de rigtige EEG-målinger, så introduceres independent component analysis (ICA) og dens estimater da dette allerede er en eksisterende og succesfuld metode for det inverse EEG-problem – for systemer med et lige antal sensorer og sources. Estimaterne fra den samlede metode sammenlignes med estimaterne fra ICA. For det lige system ses en tilstrækkelige identificering af source matrix med den samlede metode men for systemer med flere sources end sensorer fejler metoden. En alternativ test blev udført som et forsøg på

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at analysere det fundne source matrix fra et andet perspektiv. Resultatet af dette ændrer ikke den tidligere konklusion. For det praktiske perspektiv undersøges det om man kan identificere de aktive sources i forhold til det samlede antal sources igennem empiriske tests. Dog findes resultatet ikke tilstrækkelige og upålideligt.

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Introduction

The topic of this thesis arise from the increasing use of electroencephalographic measurements for a wide range of scientific purposes, especially within the medical field. By sensors placed on the head, an electroencephalography captures a mixture of electric signals caused by activity within the brain.

One essential issue concerning an electroencephalography is to recover the original source signals which was released inside the brain.

The need for source recovery is confirmed by studies showing how analysis performed on electroencephalographic measurements differs significantly from similar analysis performed directly on the original source[17]. One area of application, for which the use of the recovered source signals have shown potential, is the hearing aid industry. Here it is of interest to recover the source signals from only few sensors which potentially can be placed within a hearing aid.

Consider the issue of source recovery from a mathematical perspective. Here the electroencephalographic measurements can be modelled by a linear system of equations. From such model it is possible to recover a limited number of source signals under certain conditions. However, it is a general acknowledged issue that the true number of source signals is unknown. The task complexity of recovering the source signals from the linear system is increased in cases where the number of sources exceeds the number of sensors providing the measurements.

This thesis explores one state of the art mathematical method for source recovery, embracing the case of more sources than sensors. Overall this method, published in 2015, consist of two steps, that is receptively to recover the mixing process that the source signals have undergone and then recover the source signals. The two steps originates from two different approaches considering the mathematical orientation. The main goal of the thesis is to study the two methods with respect to proposing a united algorithm, to be applied on electroencephalographic measurements. The purpose is to support the current results of recovering source signals from electroencephalographic measurements of few sensors. Furthermore the issue of the unknown number of active source signals is considered from a perspective of practical application.

The thesis consist of a motivational part introducing electroencephalography

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and the potential use within research. Existing literature are examined, with respect to identification of state of the art approaches within source signal recovery. The Motivational part is concluded by the problem statement specifying the objective of the thesis. Next is the theoretical part. The system model is specified and the solution approach are presented based on existing methods. This is followed by an extensive study of the necessary theory. The practical aspect of the thesis includes an implementation of the proposed solution to be tested on new electroencephalographic measurements. Finally discussion and conclusion upon the achieved results are presented followed by a consideration upon further studies.

Chapter 1

Motivation

This chapter accounts for the motivation behind source extraction from an Electroencephalography (EEG). The concept of EEG is introduced along with current applications. The potential and importance of source extraction are considered and related to the hearing aid industry. The commonly applied mathematical model for EEG measurements is presented. Currently applied methods for source extraction are considered leading to a presentation of the current state of the art methods which succeeds to overcome the limitations of previous methods. Lastly, the objective of this thesis is specified.

1.1 Introduction to EEG Measurements

EEG is an imaging technique used within the medical field. EEG measures electric signals on the scalp, caused by brain activity. The human central nerve system consists of various nerve cells connecting the neurons within the brain. Nerve cells respond to certain stimuli, for instance a physical stimulus, and transmit informations between neurons. Generally speaking these activities induce local currents that are transferred throughout the nerve system. Several nearby simultaneous activations result in local potential fields, referred to as one signal source[27]. EEG measurements are provided by a number of metal electrodes, referred to as sensors, carefully placed on the human scalp. Each sensor reads the present electrical signals over time. For the source signal to reach a sensor it has to penetrate the skull, skin and several other thin layers of biological tissue. This causes an unknown distortion and reduction of a signal. It is most likely that the measurement of one sensor is a sum of multiple signals from different sources. Nor is the range of a single sensor separated from the other sensors. Thus the same signal can easily be measured by two or more sensors. The process of distortion and mixing of signals is called volume conduction [27, p. 68] [28]. From this it is clarified that EEG measurements is a mixture of fluctuating electrical signals originating from brain activities. Due to the mixing

and the nature of the signals, the true number of sources is generally considered unknown[27]. Furthermore, EEG is a subject for interfering noise. Noise signals can occur in the measurements resulting from physical movement of e.g. eyes and jawbone [30]. The concept of volume conduction is sought illustrated on figure 1.1.

The source signals are classified within four groups according to the dominant frequency. The delta wave $(0.5-4~\mathrm{Hz})$ is observed from infants and sleeping adults, the theta wave $(4-8~\mathrm{Hz})$ is observed from children and sleeping adults, the alpha wave $(8-13~\mathrm{Hz})$ is the most extensive studied brain rhythm, which is induced by an adult laying down with closed eyes. Lastly, the beta wave $(13-30~\mathrm{Hz})$ is considered the normal brain wave for adults, associated with active thinking, active attention or solving concrete problems [27, p. 11]. An example of EEG measurements within the four categories is illustrated by figure 1.2.

Generally, the distribution of EEG measurements of multiple sensors is considered multivariant Gaussian [27, p. 50]. Though the mean and covariance properties generally changes over time. Therefore, EEG measurements are considered quasistationary i.e. stationary only within small intervals. This motivates the need for segmentation of the EEG measurements to achieve signals with similar characteristics

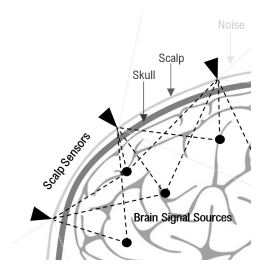


Figure 1.1: Illustration of volume conduction.



Figure 1.2: Example of time dependent EEG measurements within the four defined categories, source: [27].

1.1.1 Application

EEG performed on humans and animals have a great number of applications with both clinical and research purposes. Examples of clinical applications covers diagnosis and management of neurological disorders such as epilepsy and monitor alertness regarding coma or brain death. EEG capitalises on the procedure being non-invasive and fast. Neural activity can be measured within fractions of a second after a stimulus has been provided. These advantages contribute to the wide range of applications within research of the neural processes involved in or resulting from actions, emotions or cognition. Today such neural research are used in many different fields [30, p. 4]. The hearing aid industry is one example where this research is highly prioritised. At Eriksholm research center, which is a part of the hearing aid manufacturer Oticon, cognitive hearing science is a research area within fast development [29]. One main purpose at Eriksholm is to make it possible for a hearing aid to identify the user-intended sound source from real time EEG measurements and thereby exclude noise from elsewhere [2] [7]. It is essentially the well-known but unsolved cocktail problem which is sought improved by use of EEG. This is where EEG and occasionally so called in-ear EEG is interesting. In conjunction with the technology of beamforming, it is possible for a hearing aid to receive only signals from a specific direction.

Over the past two decades, functional integration has become an area of interest regarding EEG research [16]. Within neurobiology functional integration refers to the study of the correlation among activities in different regions of the brain. In other words, how do different parts of the brain work together to process information and conduct a response [17]. For this purpose separation and localisation of the original sources which contribute to the EEG measurement is of interest. An article from 2016 [28] points out the importance of performing analysis regarding functional integration at source level rather than at EEG level. It is argued through experiments that analysis at EEG level does not allow interpretations about the interaction between sources. This emphasises a potential for improving results within a wide range of EEG research, if the original active sources can be extracted from a specific EEG measurement.

1.1.2 Modelling

Consider the issue of extracting the activated sources from EEG measurements on the scalp. A known approach is to model the observed data by a linear system

$$y = Ax$$
.

The vector $\mathbf{y} \in \mathbb{R}^M$ is the EEG measurement of one time sample containing M sensor measurements. $\mathbf{x} \in \mathbb{R}^N$ is the corresponding N sources within the brain. The non-zero entries of \mathbf{x} represent the active sources at the time of the measurement. $\mathbf{A} \in \mathbb{R}^{M \times N}$ is an unknown transformation matrix, also referred to as the mixing matrix resembling the volume conduction. The i-th column of \mathbf{A} represents the relative projection weights from the i-th source to every sensor [6]. Representing one time sample the linear system is in general referred to as a single measurement vector model. It is only the measurement vector \mathbf{y} that is known hence it is unpossible to solve the linear system with respect to \mathbf{x} using basic linear algebra. The task, in

this case, is to identify both \mathbf{A} and then \mathbf{x} , given the measurement vector \mathbf{y} . This problem is referred to as the inverse problem of EEG. Finding \mathbf{x} from the inverse problem is referred to as source separation and localisation. Separation is to find the signal of each active source, and localisation is to place each active source signal at the right position within the source vector of dimension N, where N is the maximum number of sources.

1.1.3 Solution Method

Independent Component Analysis (ICA) is one commonly applied method to solve the inverse problem of EEG [23], [22]. ICA is a technique to find the matrix A such that the column wise elements of X is statistically independent. Thus statistical independence between the active sources is the essential assumption, which in the case of EEG are considered valid due to the volume conduction being effectively instantaneous [22, p. 3]. Application of ICA has shown great results regarding source separation of high-density EEG. However, a significant flaw to this method is that the EEG measurements are only separated into a number of sources that is equal to or less than the number of sensors [4]. Meaning that the EEG inverse problem can not be solved when it forms an under-determined system, which is the case when the maximum number of unknown sources N exceeds the number of sensors M. Such assumption undermines the reliability and usability of ICA, as the number of active sources easily exceed the number of sensors [6]. This is especially a drawback when low-density EEG are considered. Low-density EEG measurements are collected from equipment with less than 32 sensors, increasing the changes of M being less that N. However, improved capabilities of low-density EEG devices are desirable due to their relative low cost, mobility and ease to use.

This argues the importance of considering the inverse problem of EEG in the under-determined case where N > M. In the following section existing work considering the under-determined inverse problem of EEG is investigated further.

1.2 Related Work and Our Objective

As mentioned above ICA is a solid method for source separation in the case where separation into a number of sources equal to the number of sensors is adequate. The issue occurs in cases where the number of sources N exceeds the number of sensors M. To overcome this issue, an extension of ICA was suggested, referred to as the ICA mixture model [4]. Instead of identifying one overcomplete mixing matrix $\mathbf{A} \in \mathbb{R}^{M \times N}$ this approach learns N_{model} different mixing matrices $\mathbf{A}_i \in \mathbb{R}^{M \times M}$, to make computations more tractable. This method was further adapted into the Adaptive Mixture ICA (AMICA) which showed successful results regarding identification of more sources than sensors [26]. However, these successful results rely

on the assumption that no more than M out of N possible sources is simultaneously active. That is explicit that the source vector of dimension N has at most M non-zero entries. This assumption is still an essential limitation to the frame work, especially when considering low-density EEG. Other types of ICA algorithms for under-determined systems have been proposed, without overcoming the limitation of jointly active sources exceeding the number of sensors.

In 2015 O. Balkan et. al. suggested a new approach also targeting the identification of more active sources than sensors regarding EEG measurements. One method is proposed for learning \mathbf{A} from \mathbf{y} [4] and a different method is proposed for finding \mathbf{x} given \mathbf{y} and \mathbf{A} [5].

To learn \mathbf{A} the suggested method, referred to as Cov-DL, is a covariance-domain based dictionary learning algorithm. The method is based upon theory of dictionary learning and compressive sensing. Which dictates a framework for solving an underdetermined system when \mathbf{x} contains a sufficiently amount of zeros. This is similar to the constraint of ICA. However, to overcome this, the point is to transfer the EEG measurements into the covariance domain. In the covariance domain a higher dimensionality can be achieved compared to the original EEG sensor domain with dimension M. The transformation can be done when assuming a linear volume conduction and uncorrelated sources. As a result the theory of compressive sensing is found to apply to the covariance domain, allowing to learn \mathbf{A} by dictionary learning – even in the case where the active sources exceed the number of measurements.

The Cov-DL algorithm stands out from other straight forward dictionary learning methods as it does not relay on the sparsity of active sources. Where sparseness refers to the amount of non-zeros elements. This is an essential advantage when low-density EEG is considered. Cov-DL was tested and found to outperform AMICA [4]. As mentioned, the Cov-DL algorithm only learns the mixing matrix $\bf A$, resembling the volume conduction.

For the purpose of recovering \mathbf{x} , from \mathbf{y} and \mathbf{A} , a multiple measurement sparse Bayesian learning (M-SBL) algorithm is proposed. M-SBL is based on the concept of finding a set of non-zeros indices of the source vector \mathbf{x} which corresponds to finding the localisation of sources. The method builds upon the Bayesian statistic framework and it is targeting the case of more active sources than sensors. The method was proven to outperform the previously used algorithms, even when the defined recovery conditions regarding the found mixing matrix \mathbf{A} was unfulfilled [5].

One drawback, which is not fully covered in the referred literature, is that the two methods rely on the number of active sources being known. In practise this is not the case. Hence, an estimation of the number of active sources has to be considered for the algorithm to be useful in practice. To address this issue, a simple approach is to optimise the result with respect to the number active source, provided that some prior assumption of the expected result can be made.

The two state of the art methods resulting in source separation and localization will make the foundation of this thesis. Our aim is to investigate and fully understand the two methods in order to implement and test a joint algorithm – recovering the original sources \mathbf{x} from the measurements \mathbf{y} , when the number of active sources exceeds the number of measurements. Secondary it is of interest to consider the practical application of the algorithm, for instance within a hearing aid as described in section 1.1. As mentioned, the number of active sources is in general unknown in practise thus it is first of all an estimation of the number of active sources which is of interest for practical use of the algorithm. For this we want to investigate whether it is possible to estimate the number of active sources, through optimization.

Chapter 2

Problem Statement

EEG scalp measurements, a mixture of fluctuating electrical signals originating from brain activities and noise, due to distorting elements such as scalp and biological tissues, can be described as a linear system

Y = AX.

Y is the EEG scalp measurements measured from M sensors placed on the scalp, A is the mixing of the electrical signals denoted as the mixing matrix and X are the N original electrical signals, denoted as sources. Only the EEG measurements Y is known and it is of interest to identify the mixing matrix A and hereby the original sources X. The original sources have been shown significantly for practical use compared to the raw EEG scalp measurements. Especially, the under-determined case with more sources than sensors is of interest, resulting from low-density EEG devices which is beneficial due to low cost and easy application. In the linear algebraic sense, an under-determined linear system have infinitely solution provided a solution exists and is therefore difficult to solve. Two state of the art methods are seen to solve the issue with success, the covariance-domain dictionary learning (Cov-DL) and multiple sparse Bayesian learning (M-SBL) algorithms. The Cov-DL algorithm recovers the mixing matrix from the given measurements Y while the M-SBL algorithm localised and identifies the sources from the recovered mixing matrix and measurements. By combining the two state of art methods into one this could solve the inverse EEG problem – the identification of A and X given Y. However, the algorithms used the knowledges of the number of activations within the sources as this is an unknown variable in practice. Hence, a modification of the combined state of art methods is sought to increase the potential for practical use.

This motivates the following problem statement.

Based on state of the art method, how can we reproduce the recovering of original sources of brain activity from the EEG inverse problem, in the under-determined case, and how can this be modified to increase the potential of practical use such as the unknown brain activity?

From the problem statement the following sub-questions are established for clarification.

- Can we reproduce the Cov-DL algorithm to estimate a mixing matrix **A** from a over-complete EEG inverse problem with synthetic and realistic EEG scalp measurements?
- Can we reproduce the M-SBL algorithm to estimate a source matrix **X** from a over-complete EEG inverse problem with synthetic and realistic EEG scalp measurements?
- How can the number of active sources be estimated, based only on the EEG scalp measurements?

Chapter 3

System Model

Through this chapter a model representing the EEG measurements is specified. Along with the model different terminologies are introduced and described for further use in this thesis. At last the solution approach for estimating the model parameters is described, setting the outline of the remaining chapters of the thesis.

3.1 System of Linear Equations

Let $\mathbf{y} \in \mathbb{R}^M$ be some vector. By basic linear algebra \mathbf{y} can always be described as a linear combination of a coefficient matrix $\mathbf{A} \in \mathbb{R}^{M \times N}$ and some scalar vector $\mathbf{x} \in \mathbb{R}^N$ such that

$$\mathbf{y} = \mathbf{A}\mathbf{x}.\tag{3.1}$$

Let \mathbf{y} and \mathbf{A} be known, then 3.1 makes a system of M linear equations with N unknowns, referred to as a linear system.

To solve the linear system 3.1 with respect to \mathbf{x} one must look at the three different cases that can occur. The cases depend on the relation between the number of linear equations M and the number of unknowns N. For M=N, the system has one unique solution, provided that a solution exist. If the square coefficient matrix \mathbf{A} has full rank the solution can be found simply by inverting \mathbf{A} .

$$\mathbf{x} = \mathbf{A}^{-1}\mathbf{y}.$$

For M > N the system is over-determined, having more equations than unknown. There is not always a solution to an over-determined system. For M < N the system is under-determined, having fewer equations than unknowns. There exists infinitely many solutions to an under-determined system, provided that one solution exist [10, p. ix].

Consider now $\mathbf{y} \in \mathbb{R}^M$ as the observed measurements provided by M EEG sensors at time t. The linear system 3.1 is then considered as a single measurement

vector (SMV) model. Modelling the EEG measurements by the SMV model embody the following interpretations, based on chapter 1. Remember that EEG measurements basically are a mixture of original brain signals affected by volume conduction and noise. The vector \mathbf{x} is seen as the original source signals, with each entry representing the signal of one source. Thus, $\mathbf{x} \in \mathbb{R}^N$ is referred to as the source vector. N is considered the maximum number of sources, however zero entries may occur. Let k denote the number of non-zero entries in \mathbf{x} , referred to as the active sources at time t. The coefficient matrix \mathbf{A} , referred to as the mixing matrix, models the volume conduction and noise by mapping the source vector from \mathbb{R}^N to \mathbb{R}^M .

3.2 Multiple Measurement Vector Model of EEG

In practice EEG measurements are sampled over time by a certain sample frequency. Thus multiple EEG measurement vectors are achieved. Let L represent the total number of samples. The SMV model is now expanded to include L measurement vectors and noise:

$$Y = AX + E. (3.2)$$

 $\mathbf{Y} \in \mathbb{R}^{M \times L}$ is the observed measurement matrix, $\mathbf{X} \in \mathbb{R}^{N \times L}$ is the source matrix, and $\mathbf{A} \in \mathbb{R}^{M \times N}$ is the mixing matrix. Furthermore, $\mathbf{E} \in \mathbb{R}^{M \times L}$ is an additional noise matrix, to be expected from psychical measurements. The model is now referred to as the multiple measurement vector (MMV) model. As for (3.1) the solution set of the linear system (3.2) depends on the relation between N and M [10, p. 42].

As specified in chapter 1 it the case of more sources than sensors, M < N, that is of interest in this thesis.

3.2.1 Segmentation

In chapter 1 it is argued that EEG measurements are only stationary within small segments. Hence, the following segmentation is considered.

Let f be the sample frequency of the observed EEG measurement matrix \mathbf{Y} , and let t be the length of a time interval in seconds determining the duration of one segment. Here s is the segment index. As such the observed EEG measurement matrix \mathbf{Y} can be divided into stationary segments $\mathbf{Y}_s \in \mathbb{R}^{M \times L_s}$, possibly overlapping, where $L_s = tf$ is the number of samples within one segment. For each segment the MMV model (3.2) holds and is rewritten into

$$\mathbf{Y}_s = \mathbf{A}_s \mathbf{X}_s + \mathbf{E}_s, \quad \forall s. \tag{3.3}$$

Based on the assumption that each segment is stationary, it is assumed that each source signal remains either active or non-active throughout the segment. Thus, \mathbf{X}_s , consists of k non-zero rows, the active sources, and N-k zero rows, the non-active

3.3. Solution Method 15

sources. Note that the mixing matrix \mathbf{A} is not segmented in the same manner as \mathbf{Y} and \mathbf{X} , as the size of \mathbf{A} do not change relative to the number of segments. $\mathbf{A}_s \in \mathbb{R}^{M \times N}$ is the mixing matrix that corresponds to \mathbf{X}_s , \mathbf{Y}_s and \mathbf{E}_s .

In order to characterize the source matrix with respect to the number of non-zero rows, the term row sparseness is considered. Let the support supp(\mathbf{X}) denote the index set of non-zero rows of \mathbf{X} . To count the non-zero rows of a matrix the ℓ_0 -norm is defined

$$\|\mathbf{X}\|_0 \coloneqq \operatorname{card}(\operatorname{supp}(\mathbf{X})),$$

where the function card(·) gives the cardinality of the input set. The segmented source matrix \mathbf{X}_s is said to be p-sparse if it contains at most p non-zeros rows:

$$\|\mathbf{X}_s\|_0 \leq p$$
.

With respect to the model, the non-zero rows of the sources matrix make the active source signals. Define

$$k \coloneqq \|\mathbf{X}_s\|_0$$

as the number of active source signals within \mathbf{X}_s , this implies that $k \leq N$.

3.3 Solution Method

A MMV model for EEG measurements is now established. From the model the aim is to recover the source matrix \mathbf{X}_s for all segments, given only \mathbf{Y}_s . This gives an estimate of the original source signals from the brain as intended by the problem statement. In this section the solution method is presented and discussed, based on the state of the art methods lightly presented in section 1.2. This will outline the remaining chapters of the thesis.

Due to the problem statement, the case of interest is when M < N, typically resulting from low density EEG measurements. Thus, \mathbf{X} has to be recovered from an under-determined linear system. Hence, the solution must be found in the infinite solution space provided that one solution exists, thus simple linear algebra can not be used. Alternatively, numerical methods can be considered. By mathematical optimization it is possible to restrict the solution by some constraint. And then find the unique optimal solution with respect to some cost function and the corresponding constraints. The theory of compressive sensing provides a framework for solving an under-determined system when \mathbf{X} is known to have zero rows, thus being row sparse. Specifically a unique solution \mathbf{X} can be found when \mathbf{X} is M-sparse, cf. theorem A.1.1 in appendix A.1. When the mixing matrix \mathbf{A} is unknown, as in this current case, the concept of dictionary learning can be used to determine \mathbf{A} . Still under the assumption that \mathbf{X} is M-sparse.

The assumption of **X** being M-sparse corresponds to the number of active sources $k \leq M$. However, from chapter 1 it can not be justified to apply this assumption on low density EEG measurements. Hence, the theory of compressive sensing can not be applied directly on the established model, when M < N.

A method to overcome this limitation of compressive sensing, is the covariance domain dictionary learning (Cov-DL) method [4], introduced in chapter 1. The method leverages the increased dimensionality of the covariance domain in order to allow the theory of compressive sensing to apply to an under-determined system. Note that this method only applies to the process of learning \mathbf{A} , in the case where \mathbf{X} is not M-sparse. Hence, a different approach is necessary to recover \mathbf{X} .

For recovering \mathbf{X} , given both \mathbf{Y} and \mathbf{A} where M < N and $k \le N$, the method multiple sparse Bayesian learning (M-SBL)[5], introduced in chapter 1, is considered. This method takes advantage of the Bayesian statistic framework. Here, an empirical Bayesian estimation of \mathbf{X} is performed, based on the prior distribution of \mathbf{X} being defined by a data-dependent hyperparameter.

Combining the two methods allows recovery of \mathbf{A} and \mathbf{X} given low density EEG measurements \mathbf{Y} [6]. In the following two chapters each method is studied extensively, with the purpose of proposing the main algorithm in chapter 6.

Chapter 4

Covariance-Domain Dictionary Learning

Through this chapter the method covariance domain dictionary learning (Cov-DL) is presented in details. Along the presentation of the general method, necessary computational details are derived for the practical solution. The purpose is to recover the mixing matrix **A** from the MMV model, derived in chapter 3, in the underdetermined case. In the context of compressive sensing, the mixing matrix **A** in the MMV model is referred to a the dictionary matrix. That is the true mixing matrix is estimated as a dictionary matrix, through the process of dictionary learning. This will be elaborated further in the section concerning dictionary learning.

Cov-DL is an algorithm proposed by O. Balkan [4], leveraging the increased dimensionality of the covariance domain. The method has shown successful recovery of the mixing matrix \mathbf{A} , even in the non-sparse, under-determined case with more active sources k than observed measurements M, $k \geq M$. In short the algorithm consists of three steps. First the segmented MMV model of the EEG measurements is transformed into the covariance domain. Then, by the increased dimensionality of the covariance domain, it is possible to learn the transformed mixing matrix of the covariance domain, denoted by \mathbf{D} , based on the theory of compressive sensing. Here two different cases will appear dependent on the relation between the number of sources N and the found dimension of the covariance domain, which depends on the number of measurements M. Lastly, an inverse transformation is performed on the learned matrix \mathbf{D} , in order to obtain the wanted estimate of the mixing matrix \mathbf{A} . An essential aspect of this method is the prior assumption that the sources within one segment are uncorrelated, that is the rows of \mathbf{X}_s being mutually uncorrelated.

The section is inspired by the article [4] and chapter 3 in [6]. Selected general theory supporting essential parts of the method is elaborated in appendix A.

4.1 Covariance Domain Representation

Consider a single sample vector $\mathbf{y}_i \in \mathbb{R}^M$, containing EEG measurements. The covariance of \mathbf{y}_i is defined as

$$\Sigma_{\mathbf{y}_i} = \mathbb{E}[(\mathbf{y}_i - \mathbb{E}[\mathbf{y}_i])(\mathbf{y}_i - \mathbb{E}[\mathbf{y}_i])^T],$$

where $\mathbb{E}[\cdot]$ is the expected value operator. Let $\mathbf{Y}_s = [\mathbf{y}_1, \cdots, \mathbf{y}_{L_s}]$ be the observed measurements matrix containing all samples of segment s. Furthermore, assume that all sample vectors \mathbf{y}_i within one segment has zero mean and the same distribution. Then $\mathbf{Y}_s \in \mathbb{R}^{M \times L_s}$ is to be described in the covariance domain by the sample covariance $\widehat{\boldsymbol{\Sigma}}$. The sample covariance $\widehat{\boldsymbol{\Sigma}}$ is defined as the empirical covariance among the M measurements across the L_s samples. That is a $M \times M$ matrix $\widehat{\boldsymbol{\Sigma}}_{\mathbf{Y}_s} = [\sigma_{jk}]$ with entries

$$\sigma_{jk} = \frac{1}{L_s} \sum_{i=1}^{L_s} y_{ji} y_{ki}.$$

Using matrix notation the sample covariance of \mathbf{Y}_s can be written as

$$\widehat{\mathbf{\Sigma}}_{\mathbf{Y}_s} = \frac{1}{L_s} \mathbf{Y}_s \mathbf{Y}_s^T.$$

Similar, the source matrix \mathbf{X}_s can be described in the covariance domain by the sample covariance matrix:

$$\widehat{\mathbf{\Sigma}}_{\mathbf{X}_s} = \frac{1}{L_s} \mathbf{X}_s \mathbf{X}_s^T = \mathbf{\Lambda}_s + \boldsymbol{\varepsilon}.$$

The second equality comes from the assumption of the sources within \mathbf{X}_s being uncorrelated within one segment. By uncorrelated sources \mathbf{X}_s the sample covariance matrix is assumed to be nearly diagonal. Thus it can be written as $\mathbf{\Lambda}_s + \boldsymbol{\varepsilon}$ where $\mathbf{\Lambda}_s$ is a diagonal matrix consisting of the diagonal entries of $\widehat{\boldsymbol{\Sigma}}_{\mathbf{X}_s}$ and $\boldsymbol{\varepsilon}$ is a non-diagonal matrix representing the estimation error [4].

Each segment is now modelled in the covariance domain, note that the $\frac{1}{L_s}$ factor is left out in order to comply to the instruction of the source [4].

$$\mathbf{Y}_{s}\mathbf{Y}_{s}^{T} = (\mathbf{A}_{s}\mathbf{X}_{s} + \mathbf{E}_{s})(\mathbf{A}_{s}\mathbf{X}_{s} + \mathbf{E}_{s})^{T}$$

$$= (\mathbf{A}_{s}\mathbf{X}_{s})(\mathbf{A}_{s}\mathbf{X}_{s})^{T} + \mathbf{E}_{s}\mathbf{E}_{s}^{T} + \mathbf{E}_{s}(\mathbf{A}_{s}\mathbf{X}_{s})^{T} + \mathbf{A}_{s}\mathbf{X}_{s}\mathbf{E}_{s}^{T}$$

$$= \mathbf{A}_{s}\mathbf{X}_{s}\mathbf{X}_{s}^{T}\mathbf{A}_{s}^{T} + \mathbf{E}_{s}\mathbf{E}_{s}^{T} + \mathbf{E}_{s}\mathbf{X}_{s}^{T}\mathbf{A}_{s}^{T} + \mathbf{A}_{s}\mathbf{X}_{s}\mathbf{E}_{s}^{T}$$

$$= \mathbf{A}_{s}(\mathbf{\Lambda}_{s} + \varepsilon)\mathbf{A}_{s}^{T} + \mathbf{E}_{s}\mathbf{E}_{s}^{T} + \mathbf{E}_{s}\mathbf{X}_{s}^{T}\mathbf{A}_{s}^{T} + \mathbf{A}_{s}\mathbf{X}_{s}\mathbf{E}_{s}^{T}$$

$$= \mathbf{A}_{s}\mathbf{\Lambda}_{s}\mathbf{A}_{s}^{T} + \mathbf{A}_{s}\varepsilon\mathbf{A}_{s}^{T} + \mathbf{E}_{s}\mathbf{E}_{s}^{T} + \mathbf{E}_{s}\mathbf{X}_{s}^{T}\mathbf{A}_{s}^{T} + \mathbf{A}_{s}\mathbf{X}_{s}\mathbf{E}_{s}^{T}$$

$$= \mathbf{A}_{s}\mathbf{\Lambda}_{s}\mathbf{A}_{s}^{T} + \mathbf{E}_{s}\mathbf{E}_{s}^{T} + \mathbf{E}_{s}\mathbf{E}_{s}^{T} + \mathbf{E}_{s}\mathbf{X}_{s}^{T}\mathbf{A}_{s}^{T} + \mathbf{A}_{s}\mathbf{X}_{s}\mathbf{E}_{s}^{T}$$

$$= \mathbf{A}_{s}\mathbf{\Lambda}_{s}\mathbf{A}_{s}^{T} + \mathbf{E}_{s}$$

$$(4.1)$$

From (4.1) to (4.2) all terms where noise, ε and \mathbf{E}_s , is included, are aggregated in a joint noise term $\widetilde{\mathbf{E}}_s$. Next, the expression (4.2) is rewritten through a vectorization. Because the covariance matrix $\widehat{\mathbf{\Sigma}}_{\mathbf{Y}_s}$ is symmetric it is sufficient to vectorize only the lower triangular part, including the diagonal. For this purpose the function $\operatorname{vec}(\cdot)$ is defined to map a symmetric $M \times M$ matrix into a vector of size \widetilde{M} by row-wise vectorization of the lower triangular part. The increased dimension \widetilde{M} becomes

$$\widetilde{M} := \frac{M(M+1)}{2}. (4.3)$$

Furthermore, let $\operatorname{vec}^{-1}: \mathbb{R}^{\widetilde{M}} \to \mathbb{R}^{M \times M}$ be the inverse function for devectorisation.

Applying vec(·) results in the following expression, which concludes the transformation of model (3.3) into the covariance domain. Let \mathbf{a}_i be the *i*-th column of \mathbf{A}_s , as such the matrix product can be written in sum form where $\mathbf{\Lambda}_{s_{ii}}$ is the *ii*-th entry of $\mathbf{\Lambda}_s$.

$$\mathbf{Y}_{s}\mathbf{Y}_{s}^{T} = \sum_{i=1}^{N} \mathbf{a}_{i} \mathbf{\Lambda}_{s_{ii}} \mathbf{a}_{i}^{T} + \widetilde{\mathbf{E}}_{s}, \quad \mathbf{\Lambda}_{s_{ii}}$$

$$\operatorname{vec}\left(\mathbf{Y}_{s}\mathbf{Y}_{s}^{T}\right) = \sum_{i=1}^{N} \operatorname{vec}(\mathbf{a}_{i}\mathbf{a}_{i}^{T}) \mathbf{\Lambda}_{s_{ii}} + \operatorname{vec}(\widetilde{\mathbf{E}}_{s})$$

$$= \sum_{i=1}^{N} \mathbf{d}_{i} \mathbf{\Lambda}_{s_{ii}} + \operatorname{vec}(\widetilde{\mathbf{E}}_{s})$$

$$= \mathbf{D} \boldsymbol{\delta}_{s} + \operatorname{vec}(\widetilde{\mathbf{E}}_{s}), \quad \forall s.$$

$$(4.4)$$

Here $\delta_s \in \mathbb{R}^N$ contains the diagonal entries of the source sample-covariance matrix Λ_s and the matrix $\mathbf{D} \in \mathbb{R}^{\widetilde{M} \times N}$ consists of the columns $\mathbf{d}_j = \text{vec}(\mathbf{a}_j \mathbf{a}_j^T)$. Note that \mathbf{D} and δ_s are unknown while $\text{vec}(\mathbf{Y}_s \mathbf{Y}_s^T)$ is known from the observed data. By this transformation to the covariance domain, one segment is now represented by the single measurement model with \widetilde{M} "measurements".

It has been shown that this transformed model allows for identification of $k \leq \widetilde{M}$ active sources [25], which is a much weaker sparsity constraint than the original sparsity constraint $k \leq M$. The purpose of the Cov-DL algorithm is to leverage this transformed model to find the dictionary \mathbf{A} from \mathbf{D} and then still allow for $k \leq \widetilde{M}$ active sources to be recovered. That is the number of active sources are allowed to exceed the number of sensors as intended.

4.2 Recovery of the Mixing Matrix

The goal is now to learn first \mathbf{D} and then the associated mixing matrix \mathbf{A} . Two methods are considered relying on the relation between M and N. For now the noise vector is ignored.

4.2.1 Under-determined System

When $N > \widetilde{M} = \frac{M(M+1)}{2}$ the transformed model (4.4) makes an under-determined system. This is similar to the original MMV model (3.2) being under-determined when N > M. Thus, it is from the theory of compressive sensing again possible to solve the under-determined system if a certain sparsity is withhold, namely δ_s being \widetilde{M} -sparse. Assuming the sufficient sparsity on δ_s is withhold it is possible to learn the dictionary matrix of the covariance domain \mathbf{D} by traditional dictionary learning methods applied to the measurements represented in the covariance domain vec $(\mathbf{Y}_s \mathbf{Y}_s^T)$ for all segments s.

Dictionary Learning

As mentioned, within the theory of compressive sensing the matrix $\bf A$ is referred to as a dictionary matrix, as it determines how a sparse vector $\bf x$ is transformed to the original non-sparse signal. When the dictionary is not known a priori it is essential how to choose the dictionary matrix in order to achieve the best recovery, of the sparse vector $\bf x$ from measurements $\bf y$. This is clarified from the proof of theorem A.1.1 in appendix A.1. One choice is a pre-constructed dictionary. In many cases the use of a pre-constructed dictionary results in simple and fast algorithms for reconstruction of $\bf x$ [14]. However, a pre-constructed dictionary is typically fitted to a specific kind of data. For instance the discrete Fourier transform or the discrete wavelet transform are used especially for sparse representation of images [14]. Hence the results of using such dictionaries depend on how well they fit the data of interest, which is establishing a certain limitation.

The alternative option is to consider an adaptive dictionary based on a set of training data that resembles the data of interest. For this purpose learning methods are considered to empirically construct a fixed dictionary. There exist several dictionary learning algorithms. One is the K-SVD algorithm which was presented in 2006 by Elad et al. and found to outperform pre-constructed dictionaries, when computational cost is of secondary interest [1]. The concept of the K-SVD algorithm is introduced, and the more detailed algorithm is to be found in appendix A.2.

Consider, from the general MMV model (3.2), the measurement matrix $\mathbf{Y} \in \mathbb{R}^{M \times L}$ consisting of measurement vectors $\{\mathbf{y}_j\}_{j=1}^L$. Let the set of measurement vectors make a set of L training examples each forming a linear system

$$\mathbf{y}_j = \mathbf{A}\mathbf{x}_j$$
.

From the linear system one can learn a suitable dictionary $\hat{\mathbf{A}}$, and the sparse representation of the source matrix $\hat{\mathbf{X}} \in \mathbb{R}^N$ with the source vectors $\{\hat{\mathbf{x}}_j\}_{j=1}^L$. For a known sparsity constraint k the dictionary learning can be defined by the following

optimisation problem.

$$\min_{\mathbf{A}, \mathbf{X}} \sum_{j=1}^{L} \|\mathbf{y}_j - \mathbf{A}\mathbf{x}_j\|_2^2 \quad \text{subject to} \quad \|\mathbf{x}_j\|_1 \le k, \ 1 \le j \le L, \tag{4.5}$$

where both \mathbf{A} and \mathbf{x}_j are quantities to be determined. Learning the dictionary by the K-SVD algorithm consists of joint solving of the optimization problem with respect to \mathbf{A} and \mathbf{X} . An initial $\mathbf{A}_0 = [\mathbf{a}_1, \dots, \mathbf{a}_N]$ is chosen and the corresponding $\mathbf{X}_0 = [\mathbf{x}_1, \dots, \mathbf{x}_L]$ is determined, where $\mathbf{x}_j = [x_{1j}, \dots, x_{Nj}]^T$. Then, for each iteration an update rule is applied to every column of \mathbf{A}_0 , that is updating first \mathbf{a}_j for $j = 1, \dots, N$ and then the corresponding row \mathbf{x}_i where i = j. More details on the K-SVD algorithm are found in appendix A.2. The uniqueness of the $\hat{\mathbf{A}}$ depends on the recovery sparsity condition. As clarified earlier in 3.3 the recovery of a unique solution \mathbf{X}^* is only possible if k < M [6].

Application of Dictionary Learning

The establishment of an dictionary learning algorithm, K-SVD, is now used to learn the transformed dictionary matrix \mathbf{D} in (4.4). Note that \mathbf{D} corresponds to a segment s. Thus, in order to make training samples for learning \mathbf{D} a further segmentation is needed. This segmentation of \mathbf{Y}_s is indexed by s'. For convenience s will be avoided through out this chapter, as the same theory applies to all segments s. Hence, $\mathbf{Y}_{s'}$ referrers to one segment within the outer segment of measurements \mathbf{Y}_s .

The transformed and vectorized measurements $\{\text{vec}(\mathbf{Y}_{s'}\mathbf{Y}_{s'}^T), \forall s'\}$ now makes the training dataset for learning \mathbf{D} . As such each segment s' provides one training sample. Thus, the number of available training samples, denoted $L_{s'}$, depends on the chosen length of the segments. In practise this will vary with respect to the total amount of available data.

K-SVD is applied to the transformed model (4.4) and $\hat{\mathbf{D}}$ is found. Then it is possible to estimate the mixing matrix \mathbf{A} that generated \mathbf{D} through the known relation

$$\mathbf{d}_j = \text{vec}(\mathbf{a}_j \mathbf{a}_j^T).$$

For each column \mathbf{d}_j for j = 1, ..., N the following optimisation problem is solved with respect to the corresponding column \mathbf{a}_j of the mixing matrix.

$$\min_{\mathbf{a}_j} \|\mathbf{d}_j - \operatorname{vec}\left(\mathbf{a}_j \mathbf{a}_j^T\right)\|_2^2,$$

equivalent to

$$\min_{\mathbf{a}_j} \| \operatorname{vec}^{-1}(\mathbf{d}_j) - \mathbf{a}_j \mathbf{a}_j^T \|_2^2. \tag{4.6}$$

From [4] the global minimizer to (4.6) is given as $\mathbf{a}_j^* = \sqrt{\lambda_j} \mathbf{b}_j$, without further details or a source. Here λ_j is the largest eigenvalue of $\text{vec}^{-1}(\mathbf{d}_j)$,

$$\operatorname{vec}^{-1}(\mathbf{d}_{j}) = \begin{bmatrix} d_{11} & d_{12} & \cdots & d_{1N} \\ d_{21} & d_{22} & \cdots & d_{2N} \\ \vdots & \vdots & \ddots & \vdots \\ d_{N1} & d_{N2} & \cdots & d_{NN} \end{bmatrix}, \quad j = 1, \dots, N$$

evt. Redegørelse for resultatet her skal lave her and \mathbf{b}_i is the corresponding eigenvector. .

From this result each column of the mixing matrix \mathbf{A} can be estimated. Hence, it is possible to determine the mixing matrix in the case where the measurements transformed into the covariance domain makes an under-determined system. Provided however that the necessary sparsity constraint $\boldsymbol{\delta}_s$ being \widetilde{M} -sparse is withhold. Remember $\widetilde{M} \coloneqq \frac{M(M+1)}{2}$ thus M < k is allowed and the original sparsity constraint, \mathbf{X} being M-sparse, is relaxed.

4.2.2 Over-determined System

Consider again the measurements represented in the covariance domain (4.4). In the case of $\widetilde{M} > N$ an over-determined system is achieved where \mathbf{D} is high and thin. In general such a system is inconsistent. Thus, it is not possible to find \mathbf{D} by traditional dictionary learning methods and different methods most be considered. Let the set for transformed measurements be denoted by

$$\mathbf{Y}_{\text{Cov}} \coloneqq \left\{ \text{vec} \left(\mathbf{Y}_{s'} \mathbf{Y}_{s'}^T \right) \right\}_{s'=1}^{L_{s'}}.$$

When $\widetilde{M} > N$ it is expected from model (4.4) that the transformed measurements \mathbf{Y}_{Cov} live on or near a subspace of dimension N. This subspace is spanned by the columns of $\mathbf{D} \in \mathbb{R}^{\widetilde{M} \times N}$, and is denoted as $\mathcal{R}(\mathbf{D})$. To learn $\mathcal{R}(\mathbf{D})$ without having to impose any sparsity constraint on $\boldsymbol{\delta}$ it is possible to use Principal Component Analysis (PCA). The basic theory of PCA in found in appendix A.3.

PCA is applied to the set of transformed measurements \mathbf{Z} and the N first principal components are determined. The principal components form a set of basis vectors $\mathbf{U} = [\mathbf{u}_1, \dots, \mathbf{u}_N]$. That is a new basis which spans the subspace of which \mathbf{Z} lives. Thus the equality, as claimed in [4], $\mathcal{R}(\mathbf{U}) = \mathcal{R}(\mathbf{D})$ can be justified. However, this equality does not imply that $\mathbf{D} = \mathbf{U}$. In the case of two bases spanning the same vector space, namely $\mathcal{R}(\mathbf{U}) = \mathcal{R}(\mathbf{D})$, the projection operator of the given subsets must be the same. Consider the projection matrix the projection operator \mathbf{P} projecting onto the space $\mathcal{R}(\mathbf{D})$ spanned by the columns of \mathbf{D} , $\mathbf{P} : \mathbb{R}^{\widetilde{M}} \to \mathcal{R}(\mathbf{D})$. Due to \mathbf{D} having full rank it is a well known result that $\mathbf{P} = \mathbf{D}(\mathbf{D}^T\mathbf{D})^{-1}\mathbf{D}^T$. Thus $\mathcal{R}(\mathbf{U})$ and $\mathcal{R}(\mathbf{D})$ having the same projection matrix is true if and only if $\mathbf{D}(\mathbf{D}^T\mathbf{D})^{-1}\mathbf{D}^T = \mathbf{U}(\mathbf{U}^T\mathbf{U})^{-1}\mathbf{U}^T$. Now, remember from the relation between \mathbf{A} and \mathbf{D} that $\mathbf{d}_i = \mathbf{vec}(\mathbf{a}_i\mathbf{a}_i^T)$.

LÆS: bør vi bare sige at det er givet? eller helgardere os? From this it is possible to obtain **D** and then **A**, such that **D** span $\mathcal{R}(\mathbf{U})$ and $\mathbf{d}_i = \text{vec}(\mathbf{a}_i \mathbf{a}_i^T)$. This is specified by the following optimisation problem [4]

$$\min_{\{\mathbf{a}_i\}_{i=1}^N} \|\mathbf{D}(\mathbf{D}^T \mathbf{D})^{-1} \mathbf{D}^T - \mathbf{U}(\mathbf{U}^T \mathbf{U})^{-1} \mathbf{U}^T \|_F^2$$
s.t. $\mathbf{d}_i = \text{vec}(\mathbf{a}_i \mathbf{a}_i^T)$ (4.7)

where U results from PCA performed on Y_{Cov} . From the source [4] proposing the method it is only notified that that optimization problem (4.7) is minimised by quasi-Newton optimization methods. Hence the exact minimization approach can not be depicted. In the following section the optimization problem is analysed and processed in order to determine a suitable solution method.

4.2.3 Solution to Optimization Problem

The optimization problem (4.7) consists of an objective function forming a least-square problem with respect to the Frobenius norm. It is given that the squared norm, both the Euclidean and the Frobenius norm, are strictly convex [11, p.173]. Thus the objective function of (4.7) is assumed to be convex. The constraints in (4.7) is a set of quadratic equality constraints. This categorize the optimization problem as a quadratically constraint quadratic program. However, the constraint is not necessarily convex. By the constraints not being considered convex the optimization problem does not meet the requirements of a convex optimization problem. Hence the numerical solution methods for convex optimization problems, for which convergence is ensured, does not apply directly. In fact a non-convex quadratically constraint quadratic program in know to be a NP-hard problem[8]. Thus some sort of relaxation is preferred.

Due to the nature of the constraints it should be possible to reformulate the objective function to include the constraints into the objective function. That is constructing an unconstrained least-squares problem, which is a special subclass of convex optimization [9].

Let $\mathbf{D} = f(\mathbf{a}_1, \dots, \mathbf{a}_N)$, where $f(\mathbf{a}_1, \dots, \mathbf{a}_N) = \left\{ \mathbf{d}_j = \text{vec}(\mathbf{a}_j \mathbf{a}_j^T) \right\}_{j=1}^N$. Then an optimization problem without constraints is achieved and it can be solved by use of basic gradient methods, for instance the Newton method. In order to avoid an explicit expression of the inverse Hessian, which is used in the Newton method, quasi-Newton methods can be considered[3]. The general idea of quasi-Newton methods is to let the direction of search be based on a positive definite matrix which is generated from available data, with the purpose of estimating the Hessian.

Rendering of general optimization theory plus the theory of quasi-Newton methods is omitted in this thesis and the reader is referred to source [3]. For the implementation of Cov-DL in chapter 6 a predefined optimization module, using a quasi-Newton method, will be applied.

4.3 Pseudo Code of the Cov-DL Algorithm

Algorithm 1 Cov-DL

```
1: procedure Cov-DL(\mathbf{Y}_s)
              for s \leftarrow 1, \dots, L_{s'} do
 2:
                     \mathbf{y}_{\text{cov}_{s'}} = \text{vec}\left(\mathbf{Y}_{s'}\mathbf{Y}_{s'}^T\right)
 3:
 4:
              \mathbf{Y}_{\text{cov}} = \left\{ \mathbf{y}_{\text{cov}_{s'}} \right\}_{s'=1}^{L_{s'}}
 5:
 6:
              if N \ge \widetilde{M} then
 7:
                     procedure K\text{-SVD}(\mathbf{Y}_{cov})
 8:
                            returns \mathbf{D} \in \mathbb{R}^{\widetilde{M} \times N}
 9:
                     end procedure
10:
                     for j \leftarrow 1, \dots, N do
11:
                            T = \text{vec}^{-1}(\mathbf{d}_i)
12:
                             \lambda_i \leftarrow \max\{\text{eigenvalue}(\mathbf{T})\}
13:
                            \mathbf{b}_j \leftarrow \text{eigenvector}(\lambda_j)
14:
                            \mathbf{a}_{i} \leftarrow \sqrt{\lambda_{i}} \mathbf{b}_{i}
15:
                     end for
16:
                     \mathbf{A} = \{\mathbf{a}_j\}_{j=1}^N
17:
              end if
18:
19:
              if N < \widetilde{M} then
20:
                     \mathbf{procedure}\ \mathrm{PCA}(\mathbf{Y}_{\mathrm{cov}})
21:
                            returns \mathbf{U} \in \mathbb{R}^{\widetilde{M} \times N}
22:
                     end procedure
23:
                     procedure Min. A in (\|\mathbf{D}(\mathbf{D}^T\mathbf{D})^{-1}\mathbf{D}^T - \mathbf{U}(\mathbf{U}^T\mathbf{U})^{-1}\mathbf{U}^T\|_F^2)
24:
                            returns \mathbf{A} = {\{\mathbf{a}_j\}_{j=1}^N}
25:
                     end procedure
26:
27:
              end if
28: end procedure
```

4.4 Remarks

Through this chapter the theoretical aspects of the Cov-DL method proposed by [4] have been investigated in order to create algorithm 1 from which the implementation, in chapter 6, will be based. Furthermore the following remarks are considered with respect to the implementation.

The length of each time segment s has to be defined with respect to the assumption of the signals being stationary. However, it can not be assured that the

4.4. Remarks 25

stationary assumption is withhold over all segments and this will introduce an source of error. This must be taken into account in the preprocessing part for the implementation of Cov-DL when the EEG measurements are divided into segments.

For each segment a further segmentation is conducted into segments s', each serving as one sample in the covariance domain. Here the number of segments $L_{s'}$, depending on the chosen length, is most likely to influence the estimated dictionary. This is assuming that more training data will provide better results. Here a certain trade off may be considered. Longer segments s' lead to better sample covariance representation but also a fewer number of training samples within the covariance domain. Opposite, too short segments s' might compromise the sample covariance domain representation, thus the number of training sample will increase but the training samples might not be as representative. This trade off must be taken into account during the implementation of Cov-DL. Furthermore, overlapping segments might be an option for potential improvement of the Cov-DL method.

Chapter 5

Multiple Sparse Bayesian Learning

In this chapter the multiple sparse Bayesian learning (M-SBL) method is described in details, leading to an algorithm specifying the method. As the method leverage a Bayesian framework the general concept of Bayesian inference is briefly introduced prior to the M-SBL method. The chapter is generally based upon [5] where the method is applied to the MMV model, which is of interest in this thesis. Further theory is found in [31] and [32].

Consider the MMV model for a non-segmented case of EEG measurements with L samples

$$Y = AX + E. (5.1)$$

The model consist of the measurement matrix $\mathbf{Y} \in \mathbb{R}^{M \times L}$, source matrix $\mathbf{X} \in \mathbb{R}^{N \times L}$, mixing matrix $\mathbf{A} \in \mathbb{R}^{M \times N}$ and noise matrix $\mathbf{E} \in \mathbb{R}^{M \times L}$. Note that \mathbf{A} is known throughout the chapter, as it is found by COV-DL in chapter 4. The aim is to recover the source matrix \mathbf{X} in the case of fewer measurements than active sources, M < k, where $k \leq N$. In [5] it is proven that exact localization of the active sources can be achieved with M-SBL for k > M, when two sufficient conditions are satisfied. The basic approach of M-SBL is to apply Bayesian statistics to find the support set S providing the non-zero rows of the source matrix \mathbf{X} which corresponds to localization of the active sources. Finally, the value of the localized active sources can be estimated.

5.1 Bayesian Inference

The formal framework of Bayesian statistics is Bayes' theorem [20, p. 86]. The objective of Bayes' theorem is the leverage of both data and some specified prior.

This is where the distinguishes from the likelihood of classical frequentist statistics lies.

Considering now the current non-segmented MMV model (5.1) within the Bayesian framework. The model parameter – the source matrix \mathbf{X} – is wished estimated given the measurement matrix \mathbf{Y} . By Bayes' theorem the distribution of \mathbf{X} given \mathbf{Y} is constructed, that is the posterior distribution

$$p(\mathbf{X}|\mathbf{Y}) = \frac{p(\mathbf{Y}|\mathbf{X})p(\mathbf{X})}{p(\mathbf{Y})},$$

where $p(\mathbf{Y}|\mathbf{X})$ is the probability density function of \mathbf{Y} given \mathbf{X} , also referred to as the likelihood function, $p(\mathbf{X})$ is a prior distribution of \mathbf{X} and $p(\mathbf{Y})$ is the distribution of \mathbf{Y} serving as a normalizing parameter. By maximizing the posterior distribution $p(\mathbf{X}|\mathbf{Y})$ with respect to \mathbf{X} , the maximum a posteriori (MAP) estimate for the source matrix, can be found as

$$\hat{\mathbf{X}}_{\text{MAP}} = \arg \max_{\mathbf{X}} \frac{p(\mathbf{Y}|\mathbf{X})p(\mathbf{X})}{p(\mathbf{Y})}.$$

That is the estimate of $\hat{\mathbf{X}}$ with the highest posterior probability given the measurements \mathbf{Y} . In the desired case where M < N the MMV model (5.1) makes an under-determined system and potentially an infinite number of solutions exist with equal likelihoods.

Let the source matrix \mathbf{X} be seen as a variable drawn from some distribution $p(\mathbf{X})$, as such it is possible to narrow down the solution space. Assuming a prior belief that \mathbf{Y} is generated from a sparse source matrix, gives a so-called sparsity inducing prior. That is the entries of \mathbf{X} is drawn from some distribution which has a sharp, possibly infinite, spike at zero surrounded by fat tails. Here the fat tails make room for the few non-zero values, which can be seen as outliers.

For simplicity a Gaussian prior is however preferred. The use of a Gaussian distribution can almost be justified if a mixture of two Gaussian distributions are considered such that the variable is drawn from one of the two with equal likelihood. One where the variance of the distribution is close to zero, resembling the narrow spike around the mean at zero. And one with high variance resembling the fat tails.

Different MAP estimation approaches exist separated by the choice of sparsity inducing prior and optimization method. However, regardless of the approach some problems have shown to occur when using a fixed and algorithm-dependent prior. One issue occurs if the chosen prior does no assign sufficient probability to the sparse solution, leading to non-recovery. Another issue is a combinatorial number of suboptimal local solutions can occur. By use of automatic relevance determination (ARD) the problems related to the fixed sparse prior can be avoided [31, p. 20]. The main asset of this alternative approach is the use of an empirical prior. That is a flexible prior distribution depending on an unknown set of hyperparameters, which is to be learned from the data.

5.1.1 Empirical Bayesian Estimation

First assume that the likelihood function $p(\mathbf{Y}|\mathbf{X})$ is Gaussian, with noise variance $\sigma^2\mathbf{I}$. In general it is assumed that σ^2 is known. Furthermore, the noise-free case where $\sigma^2 \to 0$ will be discussed. Due to $\mathbf{y}_{\cdot j}$ consisting of measurement of individual EEG sensors, it is reasonable to assume independence. Furthermore, it is clear from the MMV model that one sample of measurements $\mathbf{y}_{\cdot j}$ only depend of one source sample $\mathbf{x}_{\cdot j}$. Hence every entry in \mathbf{Y} are assumed independently and identically distributed with likelihood

$$p(y_{ij}|x_{ij}) \sim \mathcal{N}(\mathbf{A}_{i}.\mathbf{x}_{\cdot j}, \sigma^{2})$$

$$= \frac{1}{\sigma^{2}\sqrt{2\pi}} \exp\left(-\frac{1}{2}\left(\frac{y_{ij} - \mathbf{A}_{i}.\mathbf{x}_{\cdot j}}{\sigma}\right)^{2}\right)$$

Now the empirical prior is defined, due to application of ARD, by assigning a Gaussian prior to each row in \mathbf{X} , resulting in a L-dimensional prior. Note that similar \mathbf{Y} the parameters x_{ij} are assumed to be independent and identically distributed. The empirical prior is now defined for each x_{ij} as a Gaussian distribution with zero mean and a variance controlled by an unknown hyperparameter γ_i :

$$p(x_{ij}; \gamma_i) \sim \mathcal{N}(0, \gamma_i).$$

Note that every entry of the *i*-th row is controlled by the same hyperparameter γ_i , that is one source signal over time is controlled by one hyperparameter. By combining the prior of each parameter, the prior of **X** is fully specified as follows

$$p(\mathbf{X}; \boldsymbol{\gamma}) = \prod_{i=1}^{N} p(\mathbf{x}_{i\cdot}; \gamma_i),$$

with the hyperparameter vector $\boldsymbol{\gamma} = [\gamma_1, \dots, \gamma_N]^T$. Note that the prior can be factorised over columns, resulting in

$$p(\mathbf{x}_{\cdot j}; \boldsymbol{\gamma}) = \prod_{i=1}^{N} p(x_{ij}; \gamma_i).$$

Combining the prior and the likelihood $p(\mathbf{y}_{\cdot j}|\mathbf{x}_{\cdot j})$ the posterior of the j-th column of the source matrix \mathbf{X} becomes

$$p(\mathbf{x}._{j}|\mathbf{y}._{j};\boldsymbol{\gamma}) = \frac{p(\mathbf{y}._{j}|\mathbf{x}._{j};\boldsymbol{\gamma})p(\mathbf{x}._{j};\boldsymbol{\gamma})}{p(\mathbf{y}._{j}|\boldsymbol{\gamma})}$$

$$= \frac{p(\mathbf{y}._{j}|\mathbf{x}._{j};\boldsymbol{\gamma})p(\mathbf{x}._{j};\boldsymbol{\gamma})}{\int p(\mathbf{y}._{j}|\mathbf{x}._{j})p(\mathbf{x}._{j};\boldsymbol{\gamma})} d\mathbf{x}._{j}$$

$$\approx p(\mathbf{y}._{j}|\mathbf{x}._{j};\boldsymbol{\gamma})p(\mathbf{x}._{j};\boldsymbol{\gamma})$$

$$\sim \mathcal{N}(\boldsymbol{\mu}._{j},\boldsymbol{\Sigma}), \tag{5.2}$$

where the denominator is the marginal likelihood of $\mathbf{y}_{.j}$ also referred to as the evidence. The marginalization is elaborated in the following section. Mean and covariance of (5.2) is given as.

$$\Sigma = \operatorname{Cov}(\mathbf{x}_{\cdot j}|\mathbf{y}_{\cdot j}; \boldsymbol{\gamma}) = \boldsymbol{\Gamma} - \boldsymbol{\Gamma} \mathbf{A}^T \boldsymbol{\Sigma}_y^{-1} \mathbf{A} \boldsymbol{\Gamma}, \quad \forall j = 1, \dots, L$$
 (5.3)

$$\mathcal{M} = [\boldsymbol{\mu}_{.1}, \dots, \boldsymbol{\mu}_{.L}] = \mathbb{E}[\mathbf{X}|\mathbf{Y}; \boldsymbol{\gamma}] = \mathbf{\Gamma} \mathbf{A}^T \boldsymbol{\Sigma}_y^{-1} \mathbf{Y}, \tag{5.4}$$

where $\Gamma = \operatorname{diag}(\gamma)$ and $\Sigma_y = \sigma^2 \mathbf{I} + \mathbf{A} \Gamma \mathbf{A}^T$. The derivation of the posterior mean and covariance if found in appendix B.1.

Let the posterior mean \mathcal{M} serve as the point estimate for the source matrix \mathbf{X} . It is clear that whenever $\gamma_i = 0$ the corresponding $\mathbf{x} : i$ is equal to zero with probability 1.

$$\mathbb{P}(\mathbf{x}_{i\cdot} = \mathbf{0}|\mathbf{Y}; \gamma_i = 0) = 1.$$

This ensures the posterior mean \mathcal{M} of the *i*-th row, $\boldsymbol{\mu}_{i}$, becomes zero, whenever $\gamma_{i} = 0$ as desired.

From this it is evident that for estimating the support set of \mathbf{X} it is sufficient to estimate the hyperparameter γ , from which the support set S can be extracted. Furthermore, the point estimate of \mathbf{X} , providing the source signal estimate, is given by \mathcal{M} [31, p. 147]. This leads to the actual M-SBL algorithm for which the aim is to estimate γ and the corresponding \mathcal{M} .

5.2 M-SBL for estimation of X

The M-SBL algorithm is now specified in order to estimate the hyperparameter γ and then the corresponding unknown sources \mathbf{X} . Due to the empirical Bayesian strategy the unknown source matrix \mathbf{X} is integrated out, also referred to as marginalization. By integrating the posterior with respect to the unknown sources \mathbf{X} the marginal likelihood of the observed data \mathbf{Y} is achieved [31, p. 146]

$$\mathcal{L}(\gamma; \mathbf{Y}) = \int p(\mathbf{Y}|\mathbf{X})p(\mathbf{X}; \gamma) \ d\mathbf{X}$$
$$= p(\mathbf{Y}|\gamma).$$

The resulting marginal likelihood of γ is to be maximised with respect to γ , that is the maximum likelihood estimate (MLE). From the ARD approach the MLE is considered the cost function. The $-2\log(\cdot)$ transformation is applied in order for the cost function to be minimized, and factors not depending on **Y** is removed. This

result in the following log likelihood.

$$\ell(\boldsymbol{\gamma}; \mathbf{Y}) = -2\log(p(\mathbf{Y}; \boldsymbol{\gamma}))$$

$$= -2\log\left(2\pi^{\frac{M}{2}} |\boldsymbol{\Sigma}_{y}|^{\frac{1}{2}} \exp\left(-\frac{1}{2} \sum_{j=1}^{L} \mathbf{y}_{.j}^{T} \boldsymbol{\Sigma}_{y}^{-1} \mathbf{y}_{.j}\right)\right)$$

$$= L\log(|\boldsymbol{\Sigma}_{y}|) + \sum_{j=1}^{L} \mathbf{y}_{.j}^{T} \boldsymbol{\Sigma}_{y}^{-1} \mathbf{y}_{.j}.$$
(5.5)

It is not expected that an explicit solution to the minimization problem can be found by differentiating and letting the expression equal to zero, hence it has to be solved iteratively based on a initial parameter guess $\gamma^{(0)}$. One iterative method is the expectation maximisation (EM) algorithm. In general each iteration consists of an expectation (E) step, where a function determines the expectation of the likelihood function given the currently estimated parameters. The E-step is followed by an maximization (M) step which computes the parameters by maximizing the expected likelihood found in the E-step. In this case the E-step is to compute the posterior moments using (5.3) and (5.4) while the M-step is the following update rule of γ_i [31, p.147]

$$\gamma_i^{(k+1)} = \frac{1}{L} \| \boldsymbol{\mu}_{i \cdot} \|_2^2 + \boldsymbol{\Sigma}_{ii}, \quad \forall i = 1, \dots, N.$$

The M-step is, in general, very slow on large data. An alternative is to use a fixed point update rule to fasten convergence on large data, however the resulting convergence has been found to sometimes be inferior to the convergence obtained by the above update rule[31, p.147]. The general point of a fixed-point update is to define the new value from the previous value. The fixed point updating step is here achieved by taking the derivative of the marginal log likelihood $\ell(\gamma)$ with respect to γ and equating it with zero. This leads to the following update rule which can replace the above M-step in the EM-algorithm [31, p.147]

$$\gamma_i^{(k+1)} = \frac{\frac{1}{L} \|\boldsymbol{\mu}_i\|_2^2}{1 - \gamma_i^{-1(k)} \boldsymbol{\Sigma}_{ii}}, \quad \forall i = 1, \dots, N.$$
 (5.6)

Empirically this alternative update rule have shown usefully in highly under-determined large-scale cases by driving many hyper parameters toward zero allowing for the corresponding weight in the source matrix to be discarded. For simultaneous sparse approximation problems, this is the process referred to as multiple sparse Bayesian learning, M-SBL.

From the resulting γ^* the support set S of the source matrix **X** can be extracted,

$$S = \{i | \hat{\gamma}_i \neq 0\},\$$

concluding the localization of active sources within \mathbf{X} . In practise some arbitrary small threshold can be used such that that any sufficiently small hyperparameter is discarded[31, p.149]. For identification of the active sources the estimate of the source matrix \mathbf{X} is given as $\mathbf{X}^* = \mathcal{M}^*$, with $\mathcal{M}^* = \mathbb{E}[\mathbf{X}|\mathbf{Y};\boldsymbol{\gamma}^*]$. This leads to the following estimate

$$\mathbf{X}^* = \begin{cases} \mathbf{x}_{i \cdot} = \boldsymbol{\mu}_{i \cdot}^*, & i \in S \\ \mathbf{x}_{i \cdot} = \mathbf{0}, & i \notin S \end{cases}$$

As mentioned, the case where one would like to find the noise free sparse representations should be considered. That is the limit when $\sigma^2 \to 0$. Here the M-SBL steps can be adapted easily by using a modified version the moments, given by [31, p.148] as

$$\Sigma = \left[\mathbf{I} - \mathbf{\Gamma}^{1/2} \left(\mathbf{A} \mathbf{\Gamma}^{1/2} \right)^{\dagger} \mathbf{A} \right] \mathbf{\Gamma}$$
$$\mathcal{M} = \mathbf{\Gamma}^{1/2} \left(\mathbf{A} \mathbf{\Gamma}^{1/2} \right)^{\dagger} \mathbf{Y}$$

erstat * med hat også i algoritmen where $(\cdot)^{\dagger}$ is the pseudo-inverse.

5.2.1 When k is Known

From M-SBL the number of active sources k is estimated as the number of non-zero entries in the hyperparameter γ . However, in the current scenario \mathbf{A} is estimated by Cov-DL prior to the application of M-SBL, where k is provided as input to Cov-DL, cf. chapter 4. Thus, k is known in prior to M-SBL and is therefore used as a parameter to the M-SBL algorithm. With k being known the estimation of the support set S from the non-zero rows of γ , cf. section 5.1.1, is overruled. Instead, when generating the support set S_k one choose the k largest entries of γ [5, p. 3]. The estimate of the source matrix is then found by

$$\mathbf{X}^* = \begin{cases} \mathbf{x}_{i \cdot} = \boldsymbol{\mu}_{i \cdot}^*, & i \in S_k \\ \mathbf{x}_{i \cdot} = \mathbf{0}, & i \notin S_k \end{cases}$$

Algorithm 2 M-SBL

Pseudo Code for the M-SBL Algorithm 5.2.2

1: procedure M-SBL(\mathbf{Y}, \mathbf{A}) $\gamma_{(0)} = \mathbf{1} \in \mathbb{R}^N$ 2:

```
tol = 0.0001
 3:
             while p < 3 or any(\gamma_{(p)} - \gamma_{(p-1)}) \ge tol do
  4:
                   \Gamma = \operatorname{diag}(\gamma_{(p)})
\Sigma = \Gamma - \Gamma \mathbf{A}^T \Sigma_y^{-1} \mathbf{A} \Gamma
\mathcal{M} = \Gamma \mathbf{A}^T \Sigma_y^{-1} \mathbf{Y}
  5:
  6:
  7:
                   for i = 1, ..., N do
  8:
                         \gamma_{i(p+1)} = \frac{\frac{1}{L} \|\boldsymbol{\mu}_i\|_2^2}{1 - \gamma_{i(p)}^{-1} \Sigma_{ii}}
  9:
                   end for
10:
                   p + = 1
11:
             end while
12:
             Return \mathcal{M}, \gamma
13:
14: end procedure
15: procedure Support(\mathcal{M}, \gamma, k)
             Support = \mathbf{0} \in \mathbb{R}^k
16:
             for j = 1, ..., k do
17:
                   if \gamma(\arg\max(\gamma)) != 0 then
18:
                          Support(j) = arg max(\gamma)
19:
                          \gamma(\arg\max(\gamma)) = 0
20:
                   end if
21:
             end for
22:
             \hat{\mathbf{X}} = \mathbf{0} \in \mathbb{R}^{N \times L}
23:
             for i in Support do
24:
                    \hat{\mathbf{X}}_i = \mathcal{M}_i
25:
             end for
26:
             Return \hat{\mathbf{X}}
27:
28: end procedure
```

5.2.3Sufficient Conditions for Exact Source Localization

In [5] it is proven that exact source localization is guaranteed in the under-determined case, k > M when the conditions in the following theorem are fulfilled. The theorem is based on a theoretical analysis of the minima where noise-free conditions are considered, that is letting $\sigma^2 \to 0$. Thus, it is essential that the following theorem applies to the noise less case. First, define a function $f: \mathbb{R}^{M \times N} \to \mathbb{R}^{\frac{M(M+1)}{2} \times N}$, such that for $B = f(\mathbf{A})$ the j-th column is given as $\mathbf{b}_{\cdot j} = \text{vec}(\mathbf{a}_{\cdot j}\mathbf{a}_{\cdot j}^T)$. Here the function

 $\text{vec}(\cdot)$ corresponds to the function defined in section 4.1, being a vectorization of the upper triangular part of a matrix.

Theorem 5.2.1

Given a dictionary matrix \mathbf{A} and a set of observed measurement \mathbf{Y} , M-SBL recovers the support set of any size k exactly in the noise-free case, if the following conditions are satisfied.

- 1. The active sources \mathbf{X}_S are orthogonal. That is, $\mathbf{X}_S \mathbf{X}_S^T = \mathbf{\Lambda}$, where $\mathbf{\Lambda}$ is a diagonal matrix and S the support set.
- 2. Rank $(f(\mathbf{A})) = N$.

The proof can be found in [5, p. 16].

Chapter 6

Implementation and Verification

This chapter describes the implementation process of the main algorithm, where the two algorithms Cov-DL and M-SBL from respectively chapter 4 and 5 are implemented and combined into one algorithm.

The implementation of each algorithm is initially tested on a simple simulated data set to verify the implementation. Next, both algorithms are tested on simulated data which aim to resemble real EEG measurements. By simulating the data set the true model parameters are known which allows for measuring the precision of the algorithms, based on a described error measurement. In addition different model variables are investigated in order to improve the model. Finally, the main algorithm is tested on the simulated data sets, and a conclusion is made based on the results.

6.1 Implementation of Algorithms

In this section the implementation of the main algorithm is described with use of a flowchart which illustrates the flow through code. The main algorithm consists of three main stages, an initialization, Cov-DL for recovery of \mathbf{A} and lastly M-SBL for recovery of \mathbf{X} . Considering figure 6.1, each stage of the algorithm is illustrated within one horizontal row of the flow diagram, furthermore the input and output are placed in their own row.

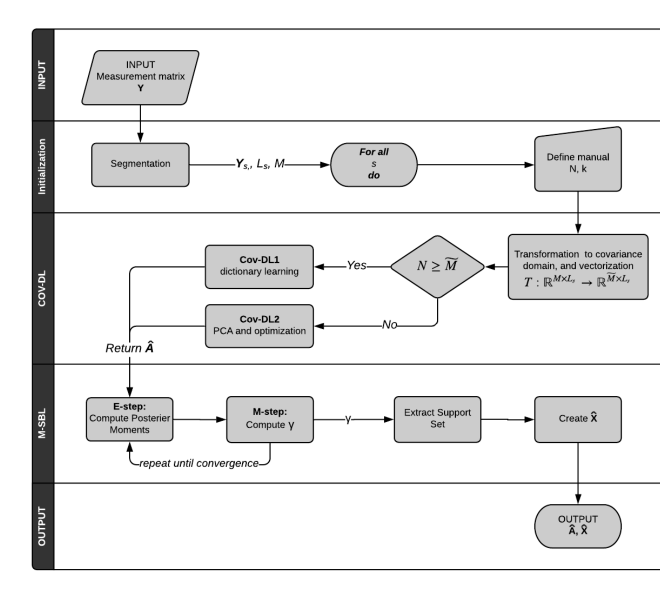


Figure 6.1: Flowchart illustrating the implementation of the main algorithm.

The input of the main algorithm consists of the measurement matrix $\mathbf{Y} \in \mathbb{R}^{M \times L}$, along with the corresponding sample frequency f. Within the initialization stage the measurement matrix \mathbf{Y} go through segmentation as described in subsection 3.2.1. The length of the segments is predefined by a time interval t in seconds such that $L_s = tf$. Each segment s is now specified by the measurement matrix $\mathbf{Y}_s \in \mathbb{R}^{M \times L_s}$. After the segmentation a loop is constructed such that for every segments s the remaining two stages, Cov-DL and M-SBL, of the main algorithm are performed. For each segment s, N and k are manually defined. This definition is either known in advance from the data or in the case of real EEG measurements they are unknown and

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a qualified guess must be made. With one segment and corresponding specifications of the expected number of active sources, the second stage of the algorithm are initialised, recovery of \mathbf{A} . The implementation of Cov-DL stage follows algorithm 1 from section 4.3 closely thus only the main steps are illustrated on the flow diagram. First the measurement matrix \mathbf{Y}_s for all segments are transformed to the covariance domain and then vectorized, resulting in an extension of the dimensionality of \mathbf{Y} from M to $\widetilde{M} = \frac{M(M+1)}{2}$. Next, the estimation of \mathbf{A} is performed from either Cov-DL1 or Cov-DL2 depending on the relation between \widetilde{M} and N, as described respectively in section 4.2.1 and 4.2.2. The estimate $\hat{\mathbf{A}}$ serves as the input to the next stage, M-SBL for recovering of \mathbf{X} , along with the measurement matrix \mathbf{Y}_s . The last stage of the main algorithm consists of the iterative EM algorithm for maximizing the marginal likelihood (5.5) with respect to γ , which is the hyperparameter from which \mathbf{X} can be determined as described in section 5.2. Lastly, the output of the main algorithm $\hat{\mathbf{X}}$ and $\hat{\mathbf{A}}$ is illustrated on the flowchart.

6.1.1 Coding Practice

The implementation of the main algorithm is performed in Python 3.6. The software and guide to run the scripts are available through appendix D.

The practical implementation process is based on module development. The established model and the three stages of the main algorithm make the system design. For each stage the necessary tasks are identified and divided into smaller modules. For each module the task is specified, and the algorithms are established and implemented. This is followed by a test of the module and possible modifications until the task is performed without error. Due to the time limitation of this project, the software was developed along side the dynamic research process. Thus, the specifications to some modules have been redefined and hence the modification process are repeated. Finally, the modules are united into one stage for which tests are performed, and lastly all the stages are united to the resulting main algorithm.

The software is based on functions, for example one module is specified by one function, for which doestrings is used, following NumPy doestring format¹ allowing insight into the structure and thoughts behind the different software elements.

For each of the stages, Cov-DL and M-SBL, verification and performance tests will be performed and described later in this chapter, followed by testing phase of the main algorithm.

6.2 Data Simulation

To test the performance of the algorithm simulated data, corresponding to the model $\mathbf{Y} = \mathbf{A}\mathbf{X}$, is needed. All data sets are simulated based on the following approach,

¹https://numpydoc.readthedocs.io/en/latest/

satisfying the sufficient conditions for recovery, displayed in theorem 5.2.1.

A source matrix $\mathbf{X} \in \mathbb{R}^{N \times L}$ is constructed, such that each row makes an independent signal which varies over L samples or a zero row. By the sources being independent the rows of \mathbf{X} becomes close to orthogonal [5], which approach the first conditions of theorem 5.2.1. Then a mixing matrix $\mathbf{A} \in \mathbb{R}^{M \times N}$ is constructed with identically distributed and independent entries. As such the source signals are randomly mixed and the mixing matrix fulfils the second condition of theorem 5.2.1. With known \mathbf{A} and \mathbf{X} the measurement matrix $\mathbf{Y} \in \mathbb{R}^{M \times L}$ is simulated according to the model, by the matrix product $\mathbf{Y} = \mathbf{A}\mathbf{X}$.

Two different kinds of data sets are simulated. One deterministic data set with simple and predictable source signals to ensure a solution and easy visualization. And a stochastic data set with randomized and fluctuating source signals to resemble realistic EEG measurements.

6.2.1 Deterministic Data Set

Two different deterministic data sets are simulated, with a different number of zero rows. One specified by $N=5,\ k=4,\ M=3$ and L=1000. That is a source matrix ${\bf X}$ with 4 independent signals and 1 zero row which is mixed into a measurement matrix with 3 measurement per sample. The second deterministic data set is specified by $N=8,\ k=4,\ M=3$ and L=1000. This is 3 additional zero rows. From the specifications the first data set comply to $N\leq \frac{M(M+1)}{2}$ which imply the use of Cov-DL2. The second data set comply to $N>\frac{M(M+1)}{2}$ and $k\leq \frac{M(M+1)}{2}$ implying the use of Cov-DL1. As such it is possible to test both branches of the Cov-DL algorithm.

The four independent source signals of X are defined by

- 1. a sinus signal $\sin(2t)$
- 2. a sawtooth signal with period $2\pi t$
- 3. a sinus signal $\sin(4t)$
- 4. a sign function of a sinus signal $\sin(3t)$

with t being a time index defined in the interval [0,4] with L samples. Each of the four signals are randomly drawn and used to construct a source matrix \mathbf{X} of size $k \times L$, then zero rows are inserted randomly, such that $\mathbf{X} \in \mathbb{R}^{N \times L}$. The mixing matrix \mathbf{A} of size $M \times N$ is randomly generated from a Gaussian distribution. By multiplying the source matrix and the mixing matrix the measurement matrix \mathbf{Y} is achieved. The deterministic data set then consist of $\{\mathbf{Y}, \mathbf{X}, \mathbf{A}\}$.

In figure 6.2 the first deterministic data set is illustrated by the source signals plotted in the top and the measurement signals plotted in the bottom. This illustrates how the source signals are transformed by the mixing matrix \mathbf{A} .

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Figure 6.2: Visualization of the signals of the source matrix **X** in comparison to the measurement signals of the measurement matrix **Y** from the deterministic data set specified by N = 5, M = 3, k = 4 and L = 1000.

6.2.2 Stochastic Data Set

The purpose of this second kind of data is to resemble EEG measurements for which the model is intended. Here different data sets are simulated depending on the chosen specifications of N, k, M and L. Every data set is constructed based on four different linear autoregressive processes of various orders, each process representing one source signal

$$- x_t^1 = \sum_{i=1}^2 \phi_i x_{t-i}^1 + w_t^1$$

$$- x_t^2 = \sum_{i=1}^2 \zeta_i x_{t-i}^2 + w_t^2$$

$$- x_t^3 = \sum_{i=1}^3 \eta_i x_{t-i}^3 + w_t^3$$

$$- x_t^4 = \sum_{i=1}^4 \xi_i x_{t-i}^4 + w_t^4$$

where ϕ, ζ, η and ξ are different model parameters and w_t^j for $j = 1, \dots, 4$ are mutually independent Gaussian distributed white noise sources. \mathbf{X} is constructed by drawing k autoregressive processes randomly each of length L among the four, if k < N zero rows are inserted randomly such that $\mathbf{X} \in \mathbb{R}^{N \times L}$. The mixing matrix \mathbf{A} of size $M \times N$ is, like the previously, generated randomly from a Gaussian distribution. By multiplying the source matrix and the mixing matrix, the measurement matrix \mathbf{Y} is achieved. The stochastic data set then consist of $\{\mathbf{Y}, \mathbf{X}, \mathbf{A}\}$.

One simulation of a stochastic data set is illustrated in figure 6.3. The illustrated data set is specified by N = 5, M = 3, k = 4 and L = 1000.

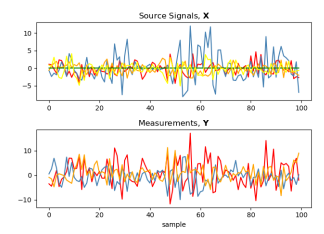


Figure 6.3: Visualization of the source signals of the source matrix **X** in comparison to the measurement signals of the measurement matrix **Y** from a stochastic data set specified by N = 5, M = 3, k = 4 and L = 1000. For simplicity only samples [0:100] are plotted.

6.2.3 Error Measurement

To evaluate performance of the algorithms, it is evident to look at the differences between the true and estimated matrices, mixing matrix \mathbf{A} and source matrix \mathbf{X} – which is possible due to the input data being simulated. For this task the mean squared error (MSE) has been chosen. The MSE measures the average squared difference between some estimated value and the true value. For $\hat{\mathbf{g}}$ being the estimate of the vector \mathbf{g} the MSE can be written as

MSE(g,
$$\hat{g}$$
) = $\frac{1}{T} \sum_{i=1}^{T} (g_i - \hat{g}_i)^2$,

with T being the number of elements in the vector \mathbf{g} .

For this project the estimates form a matrix. Here the MSE is computed for each row, which for \mathbf{X} is the estimate of one source signal, then the resulting MSE is the average over all rows. For $\mathbf{X}, \hat{\mathbf{X}} \in \mathbb{R}^{N \times L}$ the MSE is written as

$$MSE(\mathbf{X}, \mathbf{\hat{X}}) = \frac{1}{N} \sum_{i}^{N} \left(\frac{1}{L} \sum_{j=1}^{L} (\mathbf{X}_{ij} - \mathbf{\hat{X}}_{ij})^{2} \right).$$

Similarly, the MSE can be written for $\mathbf{A}, \hat{\mathbf{A}} \in \mathbb{R}^{M \times N}$.

The MSE is viewed as a measure of the quality of an estimator, in this case of how M-SBL and Cov-DL perform. The MSE considers both the variance among the estimated samples and the bias which is how far the average estimate value is from the truth[12, p.305]. Thus the larger MSE the more widely dispersed is the estimate around the true parameter. In particular, $\hat{\mathbf{X}}_1$ is considered a better estimated than $\hat{\mathbf{X}}_2$ if $\mathrm{MSE}(\mathbf{X}, \hat{\mathbf{X}}_1) < \mathrm{MSE}(\mathbf{X}, \hat{\mathbf{X}}_2)$.

6.3 Verification of Algorithms

In this section the implementation of Cov-DL and M-SBL are verified separately based on the MSE between the true and the estimated model parameters. During the tests on simulated data sets the segmentation stage is ignored by letting the simulated data form one single segment.

6.3.1 Test of Cov-DL

As seen from the flowchart 6.1 Cov-DL takes a measurement matrix \mathbf{Y} , N and k as input and return an estimation $\hat{\mathbf{A}}$ of the mixing matrix \mathbf{A} . The Cov-DL algorithm is tested on the two simulations of the deterministic data, specified in section 6.2.1.

Cov-DL1

For **Y** specified by $N > \widetilde{M}$ and $k \leq \widetilde{M}$, implying Cov-DL1, the true and estimated radius of **A** are plotted in figure 6.4 for visual comparison. Note that each matrix is vectorized such that the corresponding entries are compared. The resulting $MSE(\mathbf{A}, \hat{\mathbf{A}})$ between the true **A** and the estimated $\hat{\mathbf{A}}$ become

Tjek MSE for 0

$$MSE(A, \hat{A}) = 1.74$$

 $MSE(A, 0) = 1.40.$

From figure 6.4 it is seen that the precision of the estimate varies significant for each entry. Though, values within the a similar range are obtained. Furthermore, the $MSE(\mathbf{A}, \hat{\mathbf{A}})$ is fairly small suggesting that the estimate is acceptable. However, a smaller MSE is obtain from the estimate being a zero matrix.

Cov-DL2

For Y specified by $N \leq \widetilde{M}$, implying Cov-DL2, the true and estimated values of \mathbf{A} are plotted in figure 6.5 for visual comparison. Additionally, is the initial \mathbf{A}_{ini} , given to the optimization solver within Cov-DL2, plotted in the same figure, as standard that is a Gaussian matrix. The resulting MSE between the true \mathbf{A} and the estimated $\hat{\mathbf{A}}$ become

$$MSE(\mathbf{A}, \hat{\mathbf{A}}) = 3.00$$
$$MSE(\mathbf{A}, \mathbf{0}) = 0.90.$$

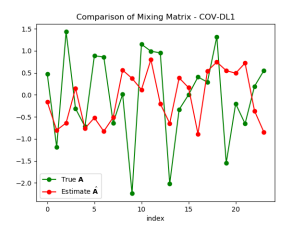
From figure 6.5 the estimate $\hat{\mathbf{A}}$ shows visual tendencies from the true \mathbf{A} . However, when it is compared to the initial guess of \mathbf{A} , \mathbf{A}_{ini} , it is observed that the estimate $\hat{\mathbf{A}}$ have moved further away from the true \mathbf{A} compared to \mathbf{A}_{ini} . This suggests some flaw within the optimization process. By printing the convergence message from the used

optimization solver, it is confirmed that the optimization process was found to be terminated successfully, with a current cost function value at 0.0 after 26 iterations. This suggests that a global minimum has been found, but the minimum, $\hat{\mathbf{A}}$, do not correspond to the true \mathbf{A} . To confirm this the following evaluations of the cost function was conducted.

$$cost(\mathbf{\hat{A}}) = 0.0$$
$$cost(\mathbf{A}_{ini}) = 1.64$$
$$cost(\mathbf{A}) = 1.65$$

These evaluations ensure that the optimization solver has managed to find the solution that minimizes the cost function. By evaluating the cost function with respect to the true $\bf A$ is it seen that it is not a global minimizer to the optimization problem. This suggests that the optimization problem, derived in section 4.2.2, do not fulfil the purpose. One reason for the optimization problem not fulfilling its purpose is that the cost function may no be convex. That is, in chapter 4 it was assumed that the cost function was convex but no further investigate regarding the truthfulness of this assumption was made. As this was not possible to do _.

Tjek dette afsnit omkring hvorfor optimisering problem ikke lykkes, cost funktion værende non-convex



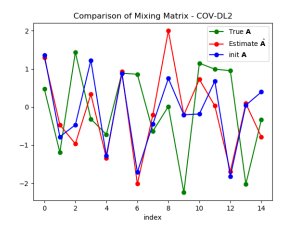


Figure 6.4: Estimated values of $\hat{\mathbf{A}}$ compared to the true values \mathbf{A}

Figure 6.5: The initial $A_{\rm ini}$ and the estimate \hat{A} compared to the true values A.

Conclusion to Estimate Â

From the results of the evaluation of cost function it is found that the estimate $\hat{\mathbf{A}}$, especially within the Cov-DL2 branch, can not be considered as a valid estimate of the mixing matrix \mathbf{A} . It is suggested that the flaw lies within either the derivation of the cost function to the optimization problem, more specifically within the assumption made throughout the derivation concerning the relation between \mathbf{A} , \mathbf{D} and \mathbf{U} . This

statement builds upon the success of the (non-documented) unit tests of the Cov-DL2 algorithm suggesting that the optimization of \mathbf{D} not depending on \mathbf{A} is possible. The appearance of this issue may suggest there is a lack within the published results [4] considering the possibility of reproducibility of the results.

Due to the time limitation of the project, the error is not investigated further, and it is concluded that the estimate of **A** is not valid hence it will not be used as an input for the next stage of the main algorithm, M-SBL. This conclusion suggests an alternative action must be considered. This is discussed further in section 6.4.

6.3.2 Test of M-SBL

From the flowchart 6.1 it seen that the M-SBL algorithm takes $\hat{\mathbf{A}}$ and \mathbf{Y} as input. The algorithm is first tested on a deterministic data set specified by M=N=k=4 and L=1000. This result will serve as a reference, showing the best possible performance due to the system having equal number of equations and unknowns, hence a unique solution exist. In order to not, let the performance of Cov-DL affect the result of M-SBL the true mixing matrix \mathbf{A} is used as input through out this section, along with the corresponding \mathbf{Y} . The resulting estimate is seen in figure 6.6. It is seen that the source signals is estimated exact, with MSE($\mathbf{X}, \hat{\mathbf{X}}$) = 0.

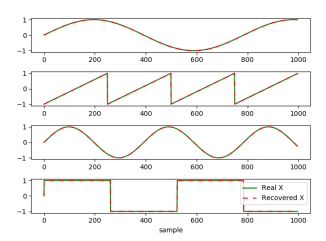


Figure 6.6: Estimated values of $\hat{\mathbf{X}}$ compared to the true values \mathbf{X} . From deterministic data \mathbf{Y} specified by $N=k=4,\ M=3$ and L=1000

Now the desired case of M < N is considered. Two tests are performed on the same two deterministic data sets, as used in the previous section, specified by M = 3, k = 4, L = 1000 and respectively N = 5 and N = 8.

The estimate $\hat{\mathbf{X}}$ is plotted in figure 6.7 and 6.8. The source signals equal to zeros of the estimate $\hat{\mathbf{X}}$ is not plotted so the figures do not visualize the exact localization of the source signals.

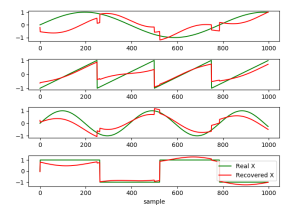


Figure 6.7: Estimated values of $\hat{\mathbf{X}}$ compared to the true values \mathbf{X} . From measurement \mathbf{Y} specified by $N=5,\ M=3,\ k=4$ and L=1000 and the true mixing matrix \mathbf{A} .

Figure 6.8: Estimated values of $\hat{\mathbf{X}}$ compared to the true values \mathbf{X} . From measurement \mathbf{Y} specified by N=8, M=3, k=4 and L=1000 and the true mixing matrix \mathbf{A} .

The resulting MSE between the true **X** and the estimate $\hat{\mathbf{X}}$ from figure 6.7 with N=5, becomes

$$MSE(\mathbf{X}, \hat{\mathbf{X}}) = 0.127.$$

From figure 6.7 it is seen that all four source signals are recovered at the right locations. As suggested by the achieved MSE the estimate is not exact, but it is clear that the estimates, by looking at figure 6.7, manage to follow the right pattern of the true signals.

The resulting MSE between the true \mathbf{X} and the estimated $\hat{\mathbf{X}}$ from figure 6.8 with N=8 thus more sparse, become

$$MSE(\mathbf{X}, \hat{\mathbf{X}}) = 0.161.$$

From figure 6.8 it is again seen that the source signals are recovered at the right location. However visually the estimates are not as. This indicates that the algorithm can manage to locate and estimate the source signals, however the increased zero rows improve the chance of dislocation and the decrease the accuracy of the estimate.

Possibilities of N = k

From the problem statement in chapter 2 it is an issue that k has to be known a priori, in order to estimate \mathbf{A} and \mathbf{X} . A short discussion in subsection 5.2.1, describes how k can be estimated within the M-SBL algorithm. However, one still needs to provide k in order to estimate \mathbf{A} , thus a qualified estimate of k can not be avoided.

Similar to k, the maximum number of active sources N is unknown in practice as it is described in chapter 1. The difference between k and N defines the number of

zero rows in \mathbf{X} . During the estimation of \mathbf{X} the localisation of the non-zero rows are, in general, significant to minimize the MSE. However, the fact that the true N can not be known for EEG measurements weakens the argument for of focusing on the localisation rather than only focusing on the value estimation of the source signals. Furthermore, it is not within the main scope of this thesis to localise the source signal. When considering the linear system, $\mathbf{Y} = \mathbf{A}\mathbf{X}$, of which the model is build upon, \mathbf{Y} does not change by removing the zero-rows of \mathbf{X} and the corresponding columns in \mathbf{A} .

From this is can be argued that letting N = k is a good estimate of N. Note in order to fulfil the sufficient conditions for the existence of a solution to the system, that k hence N is limited by \widetilde{M} .

Consider the effect of letting N=k within the M-SBL algorithm. Here it is only the estimation of the support set which is eliminated. Figure 6.9 show the estimated sources signal for a simulation of the deterministic data set now specified by N=k=4 M=3 and L=1000. The resulting MSE become

$$MSE(\mathbf{X}, \hat{\mathbf{X}}) = 0.121$$

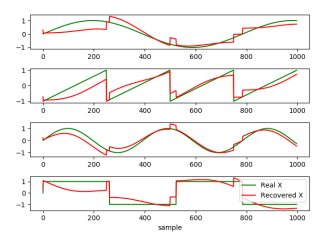


Figure 6.9: Estimated values of $\hat{\mathbf{X}}$ compared to the true values \mathbf{X} . From deterministic data \mathbf{Y} specified by $N=k=4,\ M=3$ and L=1000

From the above discussion and the results in figure 6.9 it is confirmed that letting N=k has no disadvantage with respect to the results, when the localisation of the source signal is not a priority. Thus is it chosen that N=k will be used throughout the thesis.

6.3.3 Test on Stochastic Data Sets

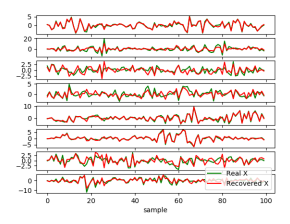
The M-SBL algorithm is now tested on two stochastic data sets which resembles the real EEG measurements. The first stochastic data set is simulated with specification

 $N=k=8,\ M=6,\ L=1000.$ The resulting estimate is plotted in figure 6.10 and the MSE becomes

$$MSE(\mathbf{X}, \hat{\mathbf{X}}) = 1.643.$$

The second stochastic data set is simulated with specification N=k=16, M=6, L=1000. This test the capabilities of the M-SBL algorithm when the distance between M and N is enlarged. The performance relative to the relation between N and M is further investigated for the main algorithm in section 6.4. The resulting estimate is plotted in figure 6.11 and the MSE becomes

$$MSE(\mathbf{X}, \hat{\mathbf{X}}) = 5.182.$$



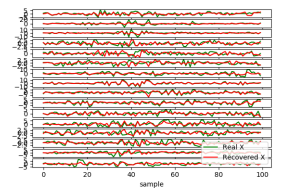


Figure 6.10: Estimated values of $\hat{\mathbf{X}}$ compared to the true values \mathbf{X} . From measurement \mathbf{Y} specified by $N=k=8,\ M=6$ and L=1000 and the true mixing matrix \mathbf{A}

Figure 6.11: Estimated values of $\hat{\mathbf{X}}$ compared to the true values \mathbf{X} . From measurement \mathbf{Y} specified by N=k=16 M=6 and L=1000 and the true mixing matrix \mathbf{A}

From figure 6.10 it is visually confirmed that the M-SBL algorithm manages to recover the source signals of the source matrix \mathbf{X} . Some source signals are nearly perfect estimated while other are having minor differences. From figure 6.11 the same tendency is seen, though more visual flaws are seen compared to figure 6.10. This result suggests that a bigger distance between M and N results in a worse performance from the M-SBL algorithm.

6.4 Test of the Main Algorithm

In this section the performance of the main algorithm is evaluated. That is the algorithm visualised in the flow diagram, figure 6.1, where the Cov-DL algorithm and the M-SBL algorithm i combined. However, as discussed due to the negative conclusion on the verification of Cov-DL an alternative has to be considered.

6.4.1 Alternative to Estimate Â

As concluded the Cov-DL algorithm does not recover a sufficient estimate of the mixing matrix \mathbf{A} , therefore a different approach is necessary.

Replacing the insufficient estimate by a fixed estimate $\hat{\mathbf{A}}_{\mathrm{fix}}$ is one immediately solution. This choice is supported by the observations from Cov-DL2 where $\mathbf{A}_{\mathrm{ini}}$ matrix provides an estimate which is happens to be a least as good as the one provided by Cov-DL. Thus, the challenge is now to determine a fixed matrix for which its characteristics resemble those of the true mixing matrix. However, from chapter 1 it is clear that no specific characteristic of the mixing matrix is known, which supports the choice of a random matrix of Gaussian distribution or similar, as it was chosen for the initial guess $\mathbf{A}_{\mathrm{init}}$ for the estimate. From this perspective three fixed mixing matrices are defined, by drawing each entry from a specified distribution:

$$\begin{aligned} \hat{\mathbf{A}}_{uni} &\sim \mathcal{U}(-1, 1) \\ \hat{\mathbf{A}}_{norm1} &\sim \mathcal{N}(0, 1) \\ \hat{\mathbf{A}}_{norm2} &\sim \mathcal{N}(0, 2) \end{aligned}$$

Note that the third matrix $\hat{\mathbf{A}}_{norm1}$ is generated the same way as the true mixing matrix of the stochastic data sets. Thus, it is expected to have the lowest MSE when compared to the true mixing matrix \mathbf{A} . However, it is of interest to investigate whether it is the best estimate of \mathbf{A} which provide the best estimate of \mathbf{X} .

A different option regarding a choice for a fixed $\hat{\mathbf{A}}$ is to utilize the ICA algorithm, described in appendix C. By the ICA algorithm it is possible to solve the EEG inverse problem for both \mathbf{A} and \mathbf{X} , in the case where $k \leq M$. Consider a simulation of a stochastic data set specified by N = k = M. Solving the system by ICA yields an estimate of \mathbf{A} . Now reduce the data set \mathbf{Y} such that $M \leq k$. Similar the estimate of \mathbf{A} is reduced by removing the same rows as in \mathbf{Y} , this yields the an estimate $\hat{\mathbf{A}}_{\text{ICA}}$ which can be used as a fixed input to M-SBL along with the corresponding reduced \mathbf{Y} .

The four different fixed estimates $\hat{\mathbf{A}}$ are tested on stochastic data sets specified by $M=10,\ N=k=16$ and L=1000, where the estimate $\hat{\mathbf{A}}_{\text{ICA}}$ has been reduced to M=10. As a reference the \mathbf{A}_{true} is included in the plot, to see the best possible $MSE(\mathbf{X},\hat{\mathbf{X}})$. To get an average performance 50 different simulations are conducted with the same specifications, each system \mathbf{X} is estimated from each of the four fixed estimates of \mathbf{A}^2 , and the MSE are computed. The resulting averaged MSE($\mathbf{A},\hat{\mathbf{A}}_{\text{fix}}$) and MSE($\mathbf{X},\hat{\mathbf{X}}$) are visualised in figure 6.12, for each of the four $\hat{\mathbf{A}}_{\text{fix}}$. Furthermore, the plotted values are found in table 6.1.

 $^{^2 \}mathrm{Note}$ that for each of the 10 repetitions four new $\mathbf{\hat{A}}_\mathrm{fix}$ are fixed.

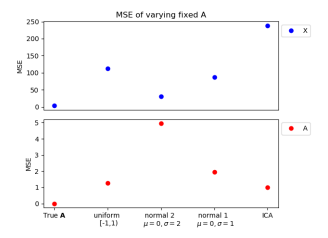


Figure 6.12: Average MSE values for each of the four fixed mixing matrix $\hat{\mathbf{A}}_{\text{fix}}$ resulting from a stochastic data set specified by M = 10, N = k = 16 and L = 1000.

	$\mathbf{\hat{A}}_{\mathrm{true}}$	$\mathbf{\hat{A}}_{ ext{uni}}$	$\mathbf{\hat{A}}_{ ext{norm2}}$	$\mathbf{\hat{A}}_{\mathrm{norm1}}$	$\hat{\mathbf{A}}_{ ext{ICA}}$
$ ext{MSE}(\mathbf{A}, \hat{\mathbf{A}}_{ ext{fix}})$	0	1.271	4.957	1.941	1.006
$\mathrm{MSE}(\mathbf{X}, \hat{\mathbf{X}})$	3.271	113.10	30.02	86.51	238.5

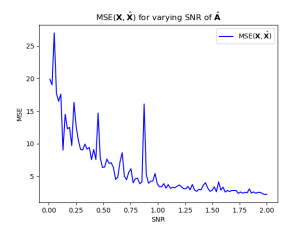
Table 6.1: Average MSE values resulting from stochastic data set specified by M = 10, N = k = 16 and L = 1000 with a fixed estimate of the mixing matrix $\hat{\mathbf{A}}_{\text{fix}}$.

From table 6.1 and figure 6.12 it is first of all seen that relation between the MSE of \mathbf{A} and \mathbf{X} is not as expected, as the lowest $\mathrm{MSE}(\mathbf{A}, \hat{\mathbf{A}}_{\mathrm{fix}})$ results in the highest $\mathrm{MSE}(\mathbf{X}, \hat{\mathbf{X}})$ and so forth. The lowest $\mathrm{MSE}(\mathbf{A}, \hat{\mathbf{A}}_{\mathrm{fix}})$ is achieved by using $\hat{\mathbf{A}}_{\mathrm{ICA}}$, which confirms that the ICA algorithm manages to estimate \mathbf{A} when $k \leq M$. However, as this do not result in the best estimate of \mathbf{X} a different choice of $\hat{\mathbf{A}}$ is still considered. The lowest $\mathrm{MSE}(\mathbf{X}, \hat{\mathbf{X}})$ is achieved by use of $\hat{\mathbf{A}}_{\mathrm{norm}}$, which resulted in the largest $\mathrm{MSE}(\mathbf{A}, \hat{\mathbf{A}}_{\mathrm{fix}})$.

As the main interest in this thesis is to identify and localize the active sources of EEG measurements a low MSE($\mathbf{X}, \hat{\mathbf{X}}$) is more desirable than a low MSE($\mathbf{A}, \hat{\mathbf{A}}_{\text{fix}}$). Furthermore, a disadvantage of using $\hat{\mathbf{A}}_{\text{ICA}}$ is the limitations in practice when k = M is not possible. From these observations a fixed estimate of the mixing matrix drawn from a normal distribution with mean 0 and variance 2, $\hat{\mathbf{A}}_{\text{norm}}$, is chosen as the alternative estimate of \mathbf{A} , and is used throughout the thesis.

Due to the unexpected relation between $MSE(\mathbf{A}, \hat{\mathbf{A}}_{fix})$ and $MSE(\mathbf{X}, \hat{\mathbf{X}})$ an additional plot is computed. Figure 6.13 show the average $MSE(\mathbf{X}, \hat{\mathbf{X}})$ as a function of $\hat{\mathbf{A}}$ with varying SNR value. This is conducted on 100 realisations of the SNR leading to the average MSE. Additionally figure 6.14 show the corresponding average MSE between the true \mathbf{A} and estimated $\hat{\mathbf{A}}$ where an increasing amount of noise is added. The SNR is also conducted on 100 realisation. Note that the noise vari-

ance is decreasing along the x-axis. The noise is generated as Gaussian white noise, with increasing variance corresponding to the desired SNR value. The SNR value is considered in the interval [0.01, 2]. From figure 6.13 it is seen that MSE decreases as the SNR decreases. This indicates as first expected that the better estimate of $\bf A$ the better estimate of $\bf X$. However, this is still a contradiction to the result seen in figure 6.12. This can however be due to the fact that the true $\bf A$ being a Gaussian matrix with $\mu=0$ and $\sigma=1$ for which Gaussian noise is added. The average MSE between the true $\bf A$ and the noisy $\hat{\bf A}$ seen in figure 6.14 is not as high as expected due to the relative large amount of noise, all being remarkably less than the corresponding MSE in table 6.1.



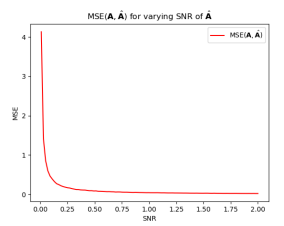


Figure 6.13: MSE($\mathbf{X}, \hat{\mathbf{X}}$) estimated from \mathbf{Y} specified by M = 6, N = k = 8 and L = 1000, as a function of SNR of given \mathbf{A} .

Figure 6.14: $MSE(\mathbf{A}, \hat{\mathbf{A}})$ where $\hat{\mathbf{A}}$ is a function of the SNR. $\hat{\mathbf{A}}$ correspond to $\hat{\mathbf{A}}$ used in figure 6.13

6.4.2 Performance Test of Main Algorithm

In order to evaluate the performance of the main algorithm, tests are conducted on several simulated stochastic data sets with different specification. The aim is to see how the relationship between N and M affect the performance, in other words how robust the algorithm is towards low density measurements. The main algorithm is tested on simulated stochastic data sets specified by M = 8, L = 1000, k = N with N in the range $N = [M+1, \ldots, 36]$, as such $k < \widetilde{M}$ is withheld insuring a solution. For each value of N ten different data sets are simulated and solved, and the average $MSE(\mathbf{X}, \hat{\mathbf{X}})$ are used as the result. The average $MSE(\mathbf{X}, \hat{\mathbf{X}})$ are plotted in figure 6.15.

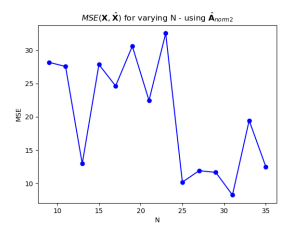


Figure 6.15: Visualization of $MSE(\mathbf{X}, \hat{\mathbf{X}})$ of the main algorithm with simulated stochastic data sets specified by M = 8, L = 1000 and k = N for $N = M + 1, \dots, 36$. Average over 10 repetitions for each N.

From figure 6.15 it is seen that the $MSE(\mathbf{X}, \hat{\mathbf{X}})$ lies in the interval [4,14], however no clear trend appears in the plot. This suggests that it is not an representative average which have been plotted, thus the test is repeated with more repetitions for value of N. The new result of the $MSE(\mathbf{X}, \hat{\mathbf{X}})$ is seen in figure 6.16.

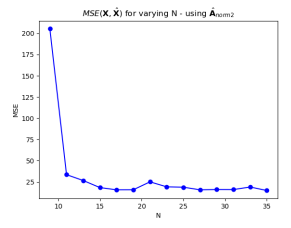


Figure 6.16: Visualization of $MSE(\mathbf{X}, \hat{\mathbf{X}})$ of the main algorithm with simulated stochastic data sets specified by M = 8, L = 1000 and k = N for $N = M + 1, \dots, 36$. Average over 500 repetitions for each N.

Figure 6.16 confirms the result of the first test. Thus, it must be that average behaviour which is seen. This suggests that the performance of the main algorithm is not affected by the relation between M and N. However, this assumption is counter intuitive and it is a contradiction to the results seen in figure 6.10 and 6.11, where

6.5. Conclusion 51

the true \mathbf{A} was utilised. Thus the choice of the alternative estimate $\hat{\mathbf{A}}_{norm}$ might have influenced the results negatively. Furthermore, it is worth to notice the relative large interval of the $\mathrm{MSE}(\mathbf{X}, \hat{\mathbf{X}})$ suggesting a vary high variance within the resulting $\mathrm{MSE}(\mathbf{X}, \hat{\mathbf{X}})$, which add a certain unreliability to the results.

6.5 Conclusion

hrough this chapter the implementation process has been described, followed by verification tests of the two main stages of the main algorithm, respectively the Cov-DL algorithm and the M-SBL algorithm.

From the test of M-SBL on stochastic data sets it was verified that the algorithm provide the expected output, and from MSE($\mathbf{X}, \hat{\mathbf{X}}$) and the corresponding visual comparison, the estimate was found to be sufficient. The verification of M-SBL was conditioned on the true mixing matrix \mathbf{A} as input, to not let the precision of the estimate $\hat{\mathbf{A}}$ from Cov-DL affect the results. Furthermore, the possibilities of letting k = N was discussed. Either N nor k is known in practise, but one has to provide the best guess for both N and k to the algorithm in order to provide corresponding number of source signals. By letting k = N one only has to guess the maximal number of active sources and not the relation between active and non-active sources, which is considered easier. Considering the consequences within the M-SBL, letting k = N will reduce the chance of dislocation, which is seen as an advantage. Furthermore, tests on the deterministic data sets confirmed that the estimated active sources were not degraded. Thus, it is confirmed that letting k = N is sufficient, and will be used when testing the main algorithm on real EEG measurements.

From the verification test of Cov-DL, providing the estimate $\hat{\mathbf{A}}$, it was found that Cov-DL did not manage to provide a sufficient result. Is been confirmed that the Cov-DL resulted in the expected output relative to the implementation, but the output did not comply with the theoretically expected result. Thus, it is concluded that the theory provided by [6] was misinterpreted, suggesting partly that the degree of reproducibility of the paper has not been sufficient. Due to the time scope of the thesis, this issue is not investigated further. However, as the estimate of \mathbf{A} resulting from Cov-DL is crucial in order to estimate the source signals from real EEG measurements, it was chosen that the best possible alternative to the original estimate must be used, in order to pursue the remaining elements of the thesis. Then, the missing estimate must be taking into account when evaluating the final results.

Different suggestions for an alternative estimate of \mathbf{A} was proposed an evaluated by the resulting $\mathrm{MSE}(\mathbf{X},\hat{\mathbf{X}})$. Here it was found that the fixed estimate $\hat{\mathbf{A}}_{\mathrm{norm}}$ generated from a normal distribution with mean 0 and variance 2 provided the best result, when tested on stochastic data sets resembling real EEG measurements.

Lastly, the performance of the main algorithm was tested on stochastic data sets. Here tests were performed on varying N in order to investigate performance

relative to the relation between M and N. For each value of N repetitions was conducted and the average $\mathrm{MSE}(\mathbf{X},\hat{\mathbf{X}})$ was evaluated. The $\mathrm{MSE}(\mathbf{X},\hat{\mathbf{X}})$ was found to lie within an interval from 2 to 25, without any characteristic trend relative to the increasing N. From this is it concluded that the performance does not rely on the relation between N and M. Despite that this was indicated by the tests where the true \mathbf{A} was utilised. Thus the lack of a precise estimate of \mathbf{A} might influence the final results.

Overall, the implementation of the main algorithm is approved. However the performance is not as good as expected. From this the main algorithm is ready to be tested on real EEG measurements in order to evaluate the performance with respect to the problem statement of this thesis. These tests are specified and conducted in the next chapter.

Chapter 7

Test on EEG measurements

The main algorithm was implemented and tested on simulated data in chapter 6. In this chapter the main algorithm is tested on EEG measurements, for which it is intended. Two different approaches are considered with respect to evaluating the resulting estimates of the source signals – ICA comparison and alpha wave analysis, respectively.

At first the provided data sets with EEG measurements are described. Followed by a test description and an analysis of the results for both of the evaluation approaches. Finally, a summary is provided to highlight the conclusions.

7.1 Data Description

For this thesis a data base of real EEG scalp measurements has been provided, from the department of electronic systems at Aalborg University. The data base consists of data sets of EEG measurements resulting from three test subjects. For each of three test subjects two data set is provided, one where the test subject sits still with open eyes and one similar but with closed eyes, resulting in a data base with 6 data sets. For the measurements an EEG cap with 32 sensors measuring the scalp EEG signal with sample frequency at 512 Hz over a period of [144,123,185,230,214,222] seconds, resulting in L = [74161,63245,94918,117900,110060,114065] samples. That is 27 channels with names and position available in EEG.chanlocs structure.

Before the data base was provided each raw data set had undergone the following preprocessing. The data were bandpass filtered between 1 and 40 Hz. Then decomposed by ICA where the independent components related to eye activity or movement was removed. Thus, for every data set 27 sensors remains. One data set then consist solely of the measurement matrix $\mathbf{Y} \in \mathbb{R}^{27 \times L}$.

Tjek lige dem måde jeg har sæt de forskellige længder ind af datasættene

7.2 Test Description

The test procedure is now described through specification of the evaluation criteria and the practical implementation of the test. Remember the aim of the developed main algorithm is to estimated the source matrix in the case where the number of sensors is less than the number of active source signals $-M < k \le N$.

7.2.1 Performance Evaluation by Comparison to ICA

From the description of ICA used on EEG measurements, cf. section 1.1.3 ICA are considered unreliable when using low-density EEG equipment where M < 32, but for $M \ge 32$ the currently considered the most reliable method for the source estimation. However note again that the true number of sources is unknown thus there is always some unreliability to the result.

From the view that the sources found by ICA where N = M is the best estimate, it is possible to let that estimate serve as a reference to be compared to the estimates achieved when M < N. In practice that is to perform ICA on a data set $\mathbf{Y} \in \mathbb{R}^{M \times L}$ resulting in $\mathbf{\hat{X}}_{ICA} \in \mathbb{R}^{N \times L}$ where M = N. Then a specific number of sensors are removed from the data set \mathbf{Y} such that M < N then and $\mathbf{\hat{X}}_{\text{Main}} \in \mathbb{R}^{N \times L}$ is estimated by the main algorithm. Then the performance of the main algorithm can be measured by comparison to the $\mathbf{\hat{X}}_{\text{ICA}}$ – does the main algorithm manage to find the same active sources as ICA, but for M < N?

In appendix C.3 the ICA algorithm is verified on simulated data without noise. It is found that ICA manage to estimate \mathbf{X} almost exact, when M=N=k. Furthermore, it is seen that for k < N ICA manage to estimate the zero rows as zero. This supports that the estimate by ICA can serve as a reference.

To compare the two estimates the MSE, cf. section 6.2.3, is used. However, an issue arises due to the fact that ICA do not manage to localise each of the found sources. That is the row of $\hat{\mathbf{X}}_{\text{ICA}}$ do not correspond to the true \mathbf{X} . Furthermore, the ICA algorithm is invariant toward the phase and the amplitude. This must necessarily worsen the resulting MSE.

The issue is cover in appendix C.3. Here a function is considered, which manage to pair and fit the rows with the lowest mutual MSE and then arrange the rows of $\hat{\mathbf{X}}_{\text{ICA}}$ such that $MSE(\mathbf{X}, \hat{\mathbf{X}}_{\text{ICA}})$ is minimised. The fitting consists of a possible phase shift and scaling of the amplitude. The right optimal fit was found through a brute-force search, however this is impossible as the possible number of combinations increases as k increases. This suggests the definition of an optimisation problem minimising the resulting MSE with respect to the combination of row indexes, possible phase and corresponding amplitude scaling. Unfortunately, a successful optimization was not achieved within the time scope of this project, thus the fitting process is not applied to the results achieved from the EEG measurements in this chapter. This factor must be taken into account when evaluating the results.

Consider again the resulting $MSE(\hat{\mathbf{X}}_{ICA}, \hat{\mathbf{X}}_{Main})$. To evaluate further on the question whether the same sources have been found a tolerance for the MSE is introduced. With $MSE(\hat{\mathbf{X}}_{ICA}, \hat{\mathbf{X}}_{Main})$ being an average over the MSE of each row within one segment a low value indicate that main part of the rows makes an estimate similar the estimate from ICA. From this perspective a tolerance for $MSE(\hat{\mathbf{X}}_{ICA}, \hat{\mathbf{X}}_{Main})$ decides whether the same sources are achieved with success. The tolerance is set to 6 due to previous observations with respect to the simulated data, especially figure 6.11 indicate that an MSE below 6 is achievable for a system where M << N with use of \mathbf{A}_{real} . It could be argued that the tolerance should be increased as the estimate of \mathbf{A} is not expected to be nearly as good, however this will could give a distorted image of the results.

7.2.2 Test Setup

The test set up is visualised in figure 7.1 by a flow diagram, showing the essential steps of the test.

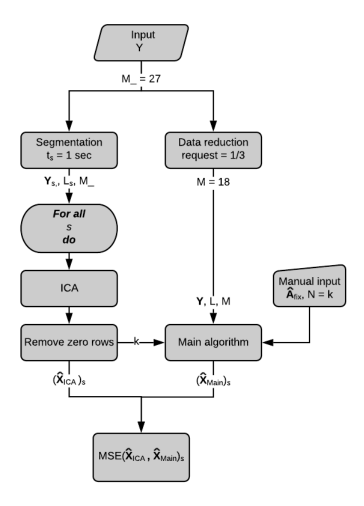


Figure 7.1: Flow diagram for visualisation of the test procedure for one data set. Example given for M < N where request = 1/3 result in M = 9.

In the flow diagram the two estimation processes are seen to run parallel but taking the same input. Prior to the application of ICA, the input is divided into segments. That is the same segmentation as inside the main algorithm, cf. section 6.1. The size of the segments is defined due to the expected stationarity of the sources. As described in the motivation chapter 1 sources are stationary if you look at sufficiently small intervals. Segments at t = 1 second is chosen from the assumption that the brain activity can be assumed stationary within short time interval. Furthermore, one must take in mind that shorter time interval lead to more segments and therefore a higher computational complexity. After the segmentation the ICA is applied to every segment, returning $\hat{\mathbf{X}}_{\text{ICA}s} \in \mathbb{R}^{M \times L_s}$. From appendix C.3 it is seen that ICA manage to estimate the non-active sources by zero rows, when no noise is present. When ICA is applied to the EEG measurements noise is expected. Thus the non-active sources defined by the average amplitude being within a tolerance interval

7.3. Results 57

around zero, defined by tol = [10E-03, -10E-03]. When the non-active sources are identified, they are removed and the resulting estimate is reduced to $\hat{\mathbf{X}}_{ICAs} \in \mathbb{R}^{k \times L_s}$. The found number of active sources k is then given as input to the main algorithm where k = N. In parallel to the ICA process the input data is reduced as specified in the previous subsection. Then the main algorithm is applied to the reduced data set. Within the main algorithm the data are like wise divided in segments and an estimate $\hat{\mathbf{X}}_{\text{Mains}} \in \mathbb{R}^{k \times L_s}$ is returned. Note that \mathbf{A}_{fix} is given as a manual input, replacing the Cov-DL algorithm as concluded in chapter 6. At the end the resulting two estimates have the same dimensions which allow for $\hat{\mathbf{X}}_{\text{Mains}}$ to be evaluated with respect to $\hat{\mathbf{X}}_{\text{ICAs}}$ by the MSE.

The described test is performed on the following three cases,

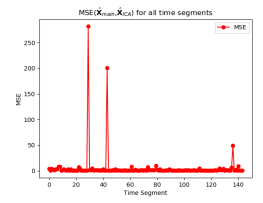
- Case 0: M = N, to see the best possible result achieved by the main algorithm.
- Case 1: M < N every third sensor is removed.
- Case 2: $M \ll N$ every second sensor is removed.

7.3 Results

For each case the test is performed on three data set — S1_Cclean, S2_Cclean and S3_Cclean. Measurements for three test subjects sitting with closed eyes in [144,185,214] seconds. The results are plotted for one data set for visual understanding, lastly the results of all three data sets are compared in a table.

7.3.1 Case 0, M = N

The results are plotted for data set S1_Cclean. The data set consist of 144 time segments with $L_s = 516$ samples and $M_{-} = 27$ sensors. Figure 7.2 show $MSE\left(\hat{\mathbf{X}}_{\text{main}}, \hat{\mathbf{X}}_{\text{ICA}}\right)$ for all segments s. ICA is applied on \mathbf{Y}_s specified by $M_{-} = 27$ and $L_s = 516$. The main algorithm is applied on \mathbf{Y}_s without any reduction hence specified by M = 27 and $L_s = 516$, given \mathbf{A}_{fix} and N = k provided from ICA. Figure 7.3 show the same plot but the y-axis is specified to the interval [-10,50] for better visualization. Furthermore is the MSE tolerance = 5 plotted, indicting for each segment whether the estimate $\hat{\mathbf{X}}_{\text{main}}$ is sufficiency close to $\hat{\mathbf{X}}_{\text{main}}$. It is seen that for a majority of the segments the MSE lies under the tolerance, but single outliers appears for which the MSE of the segment is significantly increased.



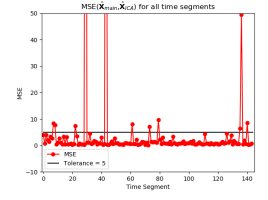
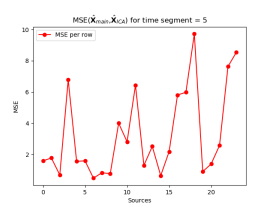
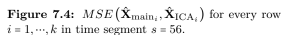


Figure 7.2: $MSE(\hat{\mathbf{X}}_{main}, \hat{\mathbf{X}}_{ICA})$ for all 144 segments

Figure 7.3: $MSE(\hat{\mathbf{X}}_{main}, \hat{\mathbf{X}}_{ICA})$ for all 144 segments. Plotted only for the y-axis interval [-10, 50] for better visualisation.

To investigate the behaviour of a single segment figure 7.4 show the MSE value computed for each row of the two estimates of a specific segment. That is $MSE\left(\hat{\mathbf{X}}_{\min_i}, \hat{\mathbf{X}}_{\text{ICA}_i}\right)$ for every row $i=1,\cdots,k$ in time segment s=56. Additionally figure ?? show and compare the corresponding estimates for four random chosen sources. This allows for visual comparison of the estimates relative to the corresponding MSE value seen in figure 7.4. Note that for better visual comparison each plotted row of $\hat{\mathbf{X}}_{ICA}$ is scaled with respect to the max value of the corresponding row in $\hat{\mathbf{X}}_{Main}$. From figure 7.4 it is seen that the estimate of each source result in a relative low MSE which indicate that the main algorithm has managed to estimate the same source as the ICA algorithm. In contradiction to this, figure 7.5 do not confirm that the estimates are close, as generally the two signals in one plot does not follow the same trend.





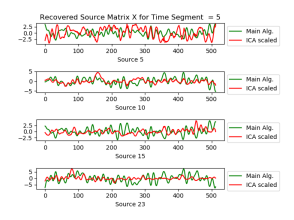


Figure 7.5: Plot of the k = 13 sources from $\hat{\mathbf{X}}_{\text{main}}$ and $\hat{\mathbf{X}}_{\text{ICA}}$ from time segment s = 56 with M = N

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The test is repeated for every data set, and the results are summarised in table 7.1. In general a low MSE is achieved in average over all segments of one data set, relative to the tolerance, and the corresponding percentage is likewise relative high, with an average at 83%. A single result is seen to deviate from the tendency, the data set of test subject 3 with closed eyes. Here a significant high average MSE value is found, indicating a majority of the segment has resulted in a significantly high MSE, while a percentage of ??? was below the tolerance. In chapter 6 it is found that the main algorithm was capable of providing an almost exact estimate for M = N when the true \mathbf{A} is provided. Thus, it is expected that the general performance is decreased in this case where the true \mathbf{A} in unknown and \mathbf{A}_{fix} is given.

The achieved results will serve as reference when analysing the results of the following cases where the main algorithm is applied on data set reduced with respect to the original data set.

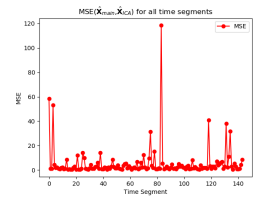
Case 0	test subject 1		test subject 2		test subject 3	
M = N	Open	Close	Open	Close	Open	Close
Average MSE()	2.913	5.172	1.572	15.06	4.753	19.44
Segments below	91	92	98	61	87	63
tolerance in $\%$	91	92	90	01	01	05

Table 7.1: Summarised results for Case 0. Test is performed on the every data set.

7.3.2 Case 1, M < N

Here the main algorithm is applied to a data set, where the number of sensors is reduced by one-third. As such ICA is applied on the original data set with segments \mathbf{Y}_s specified by $M_=27$ and $L_s=516$. The main algorithm is applied on \mathbf{Y}_s specified by M=18 and $L_s=516$, given \mathbf{A}_{fix} and N=k provided from ICA. The viewed plots correspond to those of case 0, but for reduce number of sources M < N, hence detailed plot description is omitted here.

From figure 7.6 and 7.7 it is seen that the for a majority of the segments the MSE value is close to the tolerance, but the number of outliers has increased compared to case 0, indicating the that for an increased number of segments the main algorithm do not manage to estimate enough sources sufficiently in order to stay below the tolerance.



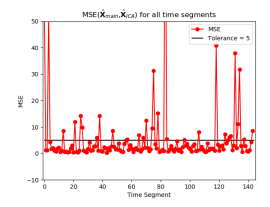
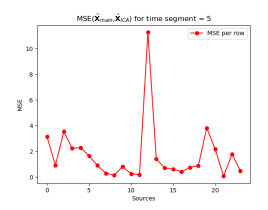


Figure 7.6: $MSE(\hat{\mathbf{X}}_{main}, \hat{\mathbf{X}}_{ICA})$ for all 144 segments

Figure 7.7: $MSE(\hat{\mathbf{X}}_{main}, \hat{\mathbf{X}}_{ICA})$ for all 144 segments. Plotted only for the y-axis interval [-10, 50] for better visualisation.

From figure 7.8 and 7.9 showing the results of segment 56, it is seen that the MSE for each source has increased slightly compared to case 0. This supports the observation from figure 7.7.



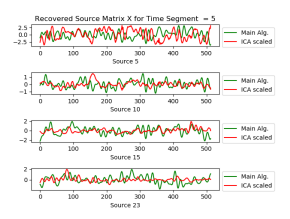


Figure 7.8: $MSE(\hat{\mathbf{X}}_{\text{main}_i}, \hat{\mathbf{X}}_{\text{ICA}_i})$ for every row $i = 1, \dots, k$ in time segment s = 56.

Figure 7.9: Plot of the k = 13 sources from $\hat{\mathbf{X}}_{\text{main}}$ and $\hat{\mathbf{X}}_{\text{ICA}}$ from time segment s = 56 with M = N

The test is repeated for every data set, and the results are summarised in table 7.2. Comparing table 7.2 to table 7.1, summarising the results of case 0, it is seen that the percentage of segments below the tolerance are decreasing, with the majority being close to 50%, thoroughly indicating that half of the time the main algorithm do not manage to provide a sufficient estimate when M = 2/3N. Furthermore both the average MSE and the corresponding percentage appears fluctuating relative to case 0 indicating some unreliability in the results.

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Case 1	test subject 1		test subject 2		test subject 3	
M < N	Open	Close	Open	Close	Open	Close
Average MSE()	9.79	5.351	13.89	15.13	6.25	18.21
Segments below tol in percent	53	80	66	46	77	48

Table 7.2: Summarised results for Case 1. Test is performed on the every data set.

7.3.3 Case 2, $M \ll N$

Here the main algorithm is applied to a data set, where the number of sensors is reduced to half. As such ICA is applied on the original data set with segments \mathbf{Y}_s specified by $M_=27$ and $L_s=516$. The main algorithm is applied on \mathbf{Y}_s specified by M=13 and $L_s=516$, given \mathbf{A}_{fix} and N=k provided from ICA. The viewed plots correspond to those of case 0 and case 1, but for further reduce number of sources $M \ll N$, hence detailed plot description is omitted.

From figure 7.10 and 7.11 it is seen that the MSE value for each segment is more widely scatted around the tolerance, compared to case 0 and 1. Outliers where the MSE value has increased significantly do also occur, similar to case 1. This indicates that the performance of the main algorithm has decreased further, compared to case 1.

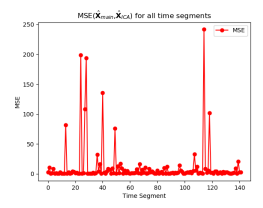


Figure 7.10: $MSE(\hat{\mathbf{X}}_{main}, \hat{\mathbf{X}}_{ICA})$ for all 144 segments

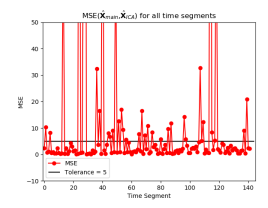
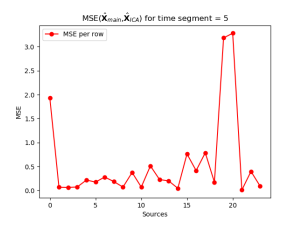


Figure 7.11: $MSE\left(\hat{\mathbf{X}}_{main}, \hat{\mathbf{X}}_{ICA}\right)$ for all 144 segments. Plotted only for the y-axis interval [-10, 50] for better visualisation.

The above indication is supported by figure 7.12 and 7.13 showing an general increase in MSE. However segment 56 still makes a fairly good example as the majority of the sources have achieves a MSE below the tolerance of 5. From figure 7.13 the increased MSE do not appear visually compared to either case 1 or case 0.



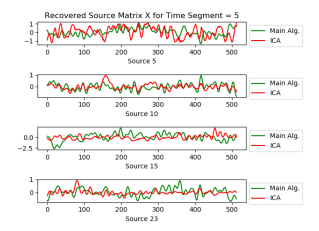


Figure 7.12: $MSE(\hat{\mathbf{X}}_{\min_i}, \hat{\mathbf{X}}_{ICA_i})$ for every row $i = 1, \dots, k$ in time segment s = 56.

Figure 7.13: Plot of the k = 13 sources from $\mathbf{\hat{X}}_{main}$ and $\mathbf{\hat{X}}_{ICA}$ from time segment s = 56 with M < N

The test is repeated for every data set, and the results are summarised in table 7.3. Comparing table 7.1 to table 7.2, summarising the results of case 1, it is generally seen that the percentage of segments below the tolerance is not decreased but improved, though without getting close to the tendency from case 0. Furthermore, the average MSE has not increased remarkably compared to case 1. As such the performance of the main algorithm in case 2 is in general not found to be worse than for case 1, however a clear improvement is not seen either.

Case 2	test subject 1		test subject 2		test subject 3	
M << N	Open	Close	Open	Close	Open	Close
Average MSE()	8.378	11.36	19.58	13.11	13.99	11.96
Segments below tol in percent	75	74	42	72	69	69

Table 7.3: Summarised results for case 2. Test is performed on the every data set.

7.3.4 Summary of Results

The main algorithm has been tested on 6 data sets of EEG measurement, for a varying relation between the number of sensors and sources – respectively case 0, 1 and 2. When the number of sensors is reduced with respect to the number of sources to be found, a significant decrease in performance was found – comparing case 0 and 1. However a corresponding decrease of performance was not found when further sensors was removed – comparing case 1 and 2.

From the conclusions made in chapter ?? it was not expected that the main algorithm would provide successful results, without estimating **A** form the data. The results of case 0 do however indicate a solid estimate provided by the main algorithm, with an average percentage of successfully estimated segments at 83%.

Furthermore, it is worth to note that the resulting MSE values has potential for improvement when considering optimization of the source localisation of the ICA estimate, cf. appendix C.3.

7.4 Alpha Wave Analysis

As mentioned in chapter 1 the source signals can be classified into four groups according to the dominant frequency [27]. It is known that when a person closes the eyes, when relaxing, the amount of alpha frequency raises and become the dominant frequency. The provided EEG measurements consist of measurements from a test subject with both open and closed eyes, hence it would be interesting to investigate the relation between the alpha frequency for open and closed eyes. The interesting part is then to compare the relation achieve from the provided measurements and the sources signals estimated by the main algorithm.

With a test of this kind, it is possible to evaluate the recovered source signals from a different perspective. Here the objective is first of all to see the behaviour with respect to the frequency, expected by the theory. Next it is interesting to investigate the aspect of analysis performed on EEG level versus analysis performed on source level, as discussed in chapter 1.

7.4.1 Test Setup

For this comparison the data sets of subject 1, S1_OClean and S1_CClean, EEG measurements of open and closed eyes respectively, will be used. It is expected that the power within the alpha frequency band is highest for the closed eyes data set, S1_CClean. To compare the amount of alpha frequency in the two datasets, a bandpass filter is used to isolate the alpha frequencies. To perform the filtering a bandpass Butterworth filter of order 5 with cut-off frequencies 8 Hz and 13 Hz will be applied to the measurements and to the source signals recovered from the main algorithm. The filtering is performed in the time domain. The filtering process is illustrated in figure 7.14. In the illustrated example only one source was investigated in both time and frequency domain, where the fast Fourier transformation (FFT) was applied [24, Chapter 9]. The source of interest was recovered from the closed-eyes dataset S1_CClean from time segment 15. The system specification used to recover the source was M = 27 and k = 14.

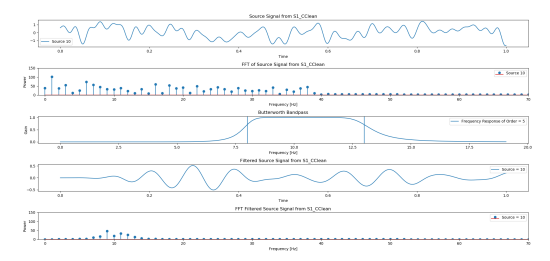


Figure 7.14: Time domain and frequency plot of a recovered source signal, filtered and non-filtered, from the time segment 15.

The first plot in figure 7.14 is the recovered source signal in the time domain. The next plot is the same source signal but transformed to the frequency domain with the FFT. The plot has been scaled to only show the frequencies from 0-70 Hz and the power from 0 to 150. The third plot illustrate the frequency response of the bandpass Butterworth filter with order 5. The vertical blue lines illustrate the cut-off frequencies at 8 Hz and 13 Hz. Plot number 4 is the recovered source signal filtered with the bandpass Butterworth filter, plotted in the time domain. The last plot is the filtered source signal plotted in the frequency domain. This verifies that the signal of interest has been filtered according to the alpha band. From the filtered source signal in the time domain, the signal resemble the alpha wave as seen in figure 1.2.

The filtering process is applied to 100 time segments of both the closed-eyes and open-eyes for respectively the raw measurement and the recovered source signals. Note that for each time segment all present sources signals or sensor measurements have been summed such that only one signal resembles each time segment.

Then for each time segment the relation between closed and open eyes is computed, with respect to power within the alpha band. The relation is defined as

$$Relation = \frac{C}{C}$$

where C is the average power from closed eyes, and O is the average power from the open eyes segment. This is done for both the raw measurements and the recovered source signals. By this it is possible to compare the relation found on source level and the relation see on EEG level.

7.4.2 Results

Figure 7.15 show an example of one time segment. To the left is the power spectrum of the filtered measurements plotted, for open and closed eyes respectively. The resulting relation between the two is 1.15. To the right is power spectrum of the filtered source signals found by the main algorithm, likewise for open and closed eyes respectively. The resulting relation between open and closed eyes is here 1.41.

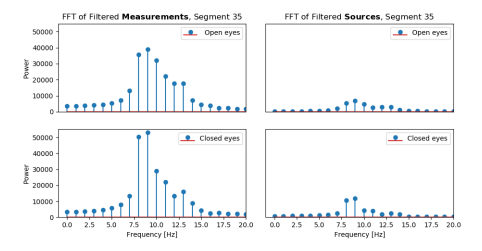


Figure 7.15: ...

By observing figure 7.15 it is seen, for the specific segment, that the power within the alpha band is significantly larger within the measurement compared to the sources. Furthermore is it seen for both the measurement and the source signals that the power has increased from open to closed eyes. Considering the calculated relations is it seen the that biggest increase in power is found on source level. This behaviour does support the theory, however the result of a single segment is not sufficient to draw any conclusion.

Figure 7.16 and 7.17 illustrate the C/O relation computed for 100 time segments, of the measurements and source signals respectively. The horizontal line in the plots marks the 1/1 relation, as such the segments where the highest power was found for closed eyes lies above the line, and opposite the segments with least power found for closed eyes lies below the line.



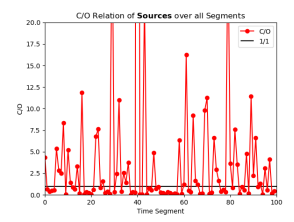


Figure 7.16: The average difference between the measurements of the open and closed eyes datasets for 100 time segments. The average difference total is 1.16.

Figure 7.17: The average difference between the recovered sources of the open and closed eyes datasets for 100 time segments. The average difference total is 2.01.

From the figures it is clear that the behaviour seen from the example of segment 35, is not a continuous behaviour. It is seen both on EEG level and source level that relation scatters round the horizontal line, indicating that the relation is not stationary over time. On figure 7.17 is it seen that the C/O relation range from near zero to beyond 20 for a few segments, indicating a significant chance in power compared to figure 7.16. With 57 out of 100 segment lying below the horizontal line it the behaviour is considered more or less random. From these observations the expected behaviour was not found. This does support the earlier findings with respect to the main algorithm, indicting a significant unreliability to the result.

With respect to the method for computing the C/O relation it could be considered whether computing the relation for every segment is the right choice. One could argue that summing the power over all segments for respectively open and closed eyes and then compute the C/O relation would yield a different result.

Chapter 8

Estimation of Active Sources

In this chapter the issue of unknown k is considered. The aim is to investigate the possibility of identifying an estimation of a non-active source signal from $\hat{\mathbf{X}}_{Main}$ when the true k is not provided to the algorithm. Instead of providing the true k one let k = N as such one ask the algorithm for N non-zero source signals, but there only k < N non-zero source signal within the synthetic data set. At first the possibilities are investigated on synthetic data, cf. section 6.2 and afterwards the real EEG data.

8.1 Empirical Test on Synthetic Data

Figure 8.1 visualise the estimate $\hat{\mathbf{X}}_{Main}$ from a stochastic data set \mathbf{Y} specified by $M=N=8,\ k=4$ and L=1000. As seen in section 6.3.3 the case of M=N should be solved almost exact by the M-SBL algorithm with true \mathbf{A} given. From the figure it is seen that the estimates of the zero rows have amplitudes close to zero, which distinguishes them from the remaining estimates, which is seen to almost exact. Due to the estimates of the zero rows being this close to zero they do not affect the MSE which is close to zero. Thus, the MSE do not indicate flaws within the estimate. Furthermore it seen that the estimates of the zero rows form a scaled copy of one of the exact estimates. These observations indicate a potential for distinguishing the estimates of zero rows and hence determine k.

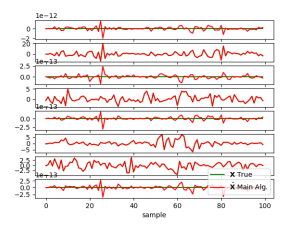


Figure 8.1: Each plot show one row of the estimate $\hat{\mathbf{X}}_{Main}$ using \mathbf{A}_{True} , compared the the corresponding true row in \mathbf{X} . The MSE is 1.196e-29. Only samples in the interval (0,100) is plotted

Consider now the desired case where M < N. Figure 8.2 visualise the estimate $\hat{\mathbf{X}}_{Main}$ from a stochastic data set \mathbf{Y} specified by M = 6, N = 8, k = 4 and L = 1000. From figure 8.2 it is seen that the estimates of the zero rows is not as close to zero as in figure 8.1. Thus, this can not be used as the indicator. However, the estimates of the zero rows still appears as a scaled replica of an estimate of a non-zero row. By a replica a signal is not considered an exact copy but a signal with similar trends over time. One attempt to locate the zero rows is to compare each row of $\hat{\mathbf{X}}_{Main}$ to every other row by the MSE, in order to check if it appears more than one time. Two rows are considered replicas if their mutual MSE is below a tolerance equal to 1. This operation is performed on the estimate plotted in figure 8.2 and gives the result displayed in table 8.1. From table 8.1 it is seen that row 2,4,6 and 8 is found to appear more than one time. Theses row indexes correspond to the zero rows of \mathbf{X} as intended. This indicates the possibility of locating the zero rows from the estimate $\hat{\mathbf{X}}_{Main}$ without providing the true k as an input.

row index	1	2	3	4	5	6	7	8
# replicas	1	3	1	2	1	4	1	3

Table 8.1: Number of replicas for each row in $\hat{\mathbf{X}}_{Main}$ of figure C.3 based on the tolerance MSE < 1.

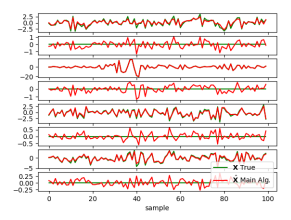


Figure 8.2: M=6, N=8, k=4, L=1000, MSE is 0.344

It is however expected that this precision must depend on the chosen tolerance for the mutual MSE. For comparison table 8.2, 8.3 and 8.4 show the result from a tolerance of 0.5, 1.5 and 2 respectively. It is observed that a tolerance of 0.5 and 2 results in a conclusion with respect to the number of zero rows – being respectively 2 and 6. From this it is clear that the tolerance is difficult to define and will affect the conclusion.

row index	1	2	3	4	5	6	7	8
# replicas	1	1	1	1	1	2	1	2

Table 8.2: Number of replicas for each row based on the tolerence MSE < 0.5.

row index	1	2	3	4	5	6	7	8
# replicas	1	4	1	3	1	4	1	3

Table 8.3: Number of replicas for each row based on the tolerence MSE < 1.5.

row index	1	2	3	4	5	6	7	8
# replicas	2	6	1	4	2	4	1	4

Table 8.4: Number of replicas for each row based on the tolerence MSE < 2.

The results so far have relied on the true \mathbf{A} as an input the main algorithm, due to the conclusion of chapter 6 where the estimate of \mathbf{A} is abandoned. Thus, this gives the results to be expected conditioned on an exact estimate of \mathbf{A} which this thesis does not manage to provide.

Now the investigations are repeated but with use of the main algorithm for which a fixed \mathbf{A} is provided as input, cf. 6.4. Similar to figure 8.2, figure 8.3 show the estimates $\hat{\mathbf{X}}_{Main}$ compared to the true \mathbf{X} . As expected, according to the results from section 6.4.2, it is generally seen from figure 8.3 that every row of the estimate is less accurate as a result of using a fixed \mathbf{A} instead of the true \mathbf{A} . Table 8.5 shows the replica count with an MSE tolerance at 1. From the table it would be concluded that 7 out of the 8 rows are zero rows, while the true number is 4. This could indicate that the tolerance is set to high. Table 8.6 show the replica count for an MSE tolerance at 0.5. From table 8.6 it is seen that the number of replicas is reduced, however, it still does not result in the right conclusion.

row index	1	2	3	4	5	6	7	8
# replicas	3	5	2	2	4	1	4	5

Table 8.5: Number of replicas for each row based on the tolerence MSE < 1.

row index	1	2	3	4	5	6	7	8
# replicas	2	2	2	2	1	1	1	3

Table 8.6: Number of replicas for each row based on the tolerance MSE < 0.5.

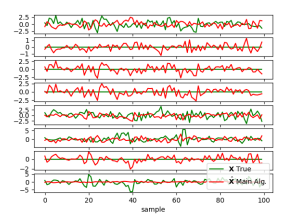


Figure 8.3: M = 6, N = 8, k = 4, MSE is 128.7

From the observations made through this investigation, based on synthetic data, the following conclusions are made. From figure 8.2 and table 6.1 a potential is found with respect to identifying the zero rows within the estimate. Here the zero rows are identified as the rows of the estimate for which similar signals appear in other rows indicating that no new estimate has been computed. For figure ?? and table

8.5 a fixed **A** is used in the main algorithm, as it will be when applied to real EEG data. Here it has not been possible to identify the zero rows correctly, based on the replica count. Thus, it must be concluded that the method is not reliable when the estimated is computed by the developed main algorithm. However, it is essential that a potential was found under ideal conditions, due to the results from the main algorithm being on reliable as concluded in chapter 6.

However, to finish the investigation, the method replica count has been applied to the estimation of real EEG data. This is done due to the possibility of seen a different behaviour from the real EEG data compared to the synthetic data.

8.2 Real EEG Measurements

Let's look at estimation of active sources, k, on the real EEG measurements. For this estimation one can not compare the estimation to the real sources as in the previous section. Instead, one applies the estimation tool, looking for replicas, on the real EEG measurements and makes a conclusion from the observed result.

For the estimation one will only look at one time segment to verify that the estimation of active sources can be applied on real EEG measurements. The estimation will be performed on the S1_Cclean EEG data set with $\frac{1}{2}M < N$ – every second sensor is removed. The specification on this data set are M=13 and the time segment of interest is s=10. For the estimation one must give an initial guess for the number of active sources. Remember that k>M and k< N. k=17 has been chosen as a choice for how many active sources exists in time segment s=10.

Figure 8.4 visualize the recovered source matrix $\hat{\mathbf{X}}$ from time segment s=10 for M=13 and k=17.

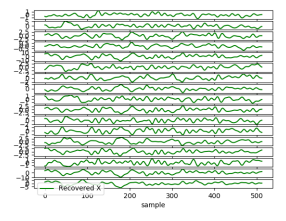


Figure 8.4: The recovered source matrix $\hat{\mathbf{X}}$ with k = 17 active source for the EEG data set with M = 13.

From figure 8.4 one see that all k active sources are visible and there seem not to be any visible indication of a active source being non-active.

From the list of replicates from figure 8.4 only four sources are the 'real' active source while the rest are replicates of those four. The 'real' active sources can be found in row 2, 5, 16 and 17. This leads to the conclusion that for time segment s=10 with system specification M=13 for the S1_Cclean EEG data set only have k=4 active sources.

Chapter 9

Discussion

The purpose of this thesis was to investigate the possibility of reproducing state of the art methods and results for localization and identification of source signals, from low-density scalp EEG measurements inducing an under-determined system. The considered state of the art methods is multiple sparse Bayesian learning (M-SBL)[5] and covariance-domain dictionary learning (Cov-DL)[4], both published by O. Balkan, et al. in the year 2014 and 2015 respectively. It was found that this task was not easy as they dit not provide any software to reproduce their results. The resulting combination of Cov-DL and M-SBL, the main algorithm, did not manage to solve the inverse EEG problem of recovering the mixing matrix A and a source matrix X successfully, from the EEG measurements. From the verification of the Cov-DL method, it was concluded that it failed to provide a sufficient estimate Â, when applied on simulated data. Due to not having a successful estimate of A the recovery of the X is compromised when using the M-SBL method. However, when using the true **A**, the M-SBL provide an estimate $\hat{\mathbf{X}}$ sufficiently close to the real source matrix, in the under-determined case M < N. In the main algorithm the estimate $\hat{\mathbf{A}}$ from Cov-DL was replaced by a fixed mixing matrix \mathbf{A}_{fix} . The final performance of the main algorithm was found very significantly, hence a sufficient performance was not confirmed. As expected a similar conclusion was to be drawn from tests on real EEG measurement.

In chapter 6 it was concluded that the implementation of the Cov-DL method is unsuccessful. The issue was located to the definition of the optimization problem determining the columns of $\hat{\mathbf{A}}$. This does question the reproducibility of the scientific article [4] which has been used as the main source. The article [4] did not provide any code or implementation specifications. Likewise, it was not possible to recreate or access the exact data, which was used to provide the results presented in the article. Thus, the intention was never to recreate the exact results from the article but rather to demonstrate the conclusion, that the method manage to provide results of a certain success rate. One could argue that testing the implementation on the

same data would lead to a different conclusion. However, this was sought approached by the stochastic simulated data, cf. section 6.2, which was created with inspiration from the article.

In chapter 6 it was concluded that the M-SBL method manage to successfully estimate \mathbf{X} when applied to the simulated stochastic data sets and given the true \mathbf{A} . Though, the performance was found to decrease slightly as the number of sources increases relative to the number of sensors. Regarding the reproducibility of the article [5] the results indicates the provided information about M-SBL has been sufficient. However, this article did, as [4], not provide any code or data disabling the possibility of recreating the exact results. In [5] tests were conducted on random simulations of \mathbf{A} and \mathbf{X} with various noise level added. Thus, due to no counter arguments, the tests of M-SBL in this thesis were conducted on the simulated data sets already created for the tests of Cov-DL, with inspiration from [5]. With respect to the main algorithm, uniting Cov-DL and M-SBL, an alternative to the estimate of \mathbf{A} was necessary. Through empirical tests, which is discussed later, a \mathbf{A}_{fix} was chosen to replace the estimate from Cov-DL. With \mathbf{A}_{fix} the main algorithm manage to estimate \mathbf{X} but with a significant higher error compared to the use of the true \mathbf{A} , which is not considered to be successful.

The performance test of the main algorithm was first conducted on simulated stochastic data resembling real EEG measurements. Inspired by [4] the sources was synthesised by independent auto-regressive processes. The true **A** however, was simply generated by a Gaussian distribution. This choice was based on a lack of information to point in different directions. Instead of the true **A** being chosen as a stochastic matrix a deterministic matrix could have been chosen instead. The choice of the stochastic true **A** of the simulated data sets could affect the results when testing the fixed alternatives for the mixing matrix estimated by Cov-DL. From the test, cf. section 6.4 it was found that a Gaussian fixed **A**, generated with same specifications as the true **A**, did not lead to the best recovery of the **X**, which went against the natural expectation – the better estimate of **A** the better estimate of **X**. Here a fixed **A** with Gaussian distribution of higher variance provided the best estimate of **X**. It is here one could argue that the stochastic true **A** had an influence to the results. Furthermore, the choice of error measurement might also be reflected in the results. Not being sufficient towards the purpose.

The common choice of error measurement throughout the thesis was the mean-square error (MSE). The MSE measures the performance of an estimate with respect to the true value, by comparing each element and summing the squared error. Hence, the MSE is providing a measure of how far the estimate is from the true value. This was considered a sufficient error measurement for evaluation of Cov-DL and M-SBL. However, one challenge when using MSE is to set a tolerance defining when an estimate is considered successful. It could be argued that set tolerance should vary with respect to the data of interest. Though, it was not chosen to evaluate to the

algorithms with respect to success rate. By this the above challenge was avoided, and the method replaced by a more soft evaluation based on the MSE. A different choice of error measurement could have been the use of correlation between variables of the estimate and the ground truth. With respect to the comparison of the main algorithm to ICA as the ground truth, to be discussed next, using the correlations as the evaluation might have overcome the scaling issue with respect to ICA.

Consider now the performance test of the main algorithm on real EEG measurements. The choice of evaluating the performance by considering the ICA solution as the ground truth has been crucial. First of all the foundation for the evaluation was not ideal. It is an issue that the performance of the main algorithm on the simulated data was not as good expected, presumably due to the estimate of A been replaced by a fixed matrix. Thus, it is not reasonable to trust the results when the algorithm is applied to data for which the true results are not known at all. However, it can be argued that comparing the obtained result to the best-known solution, in this case provided by ICA, a small error will indicated an acceptable performance from the main algorithm. The arguments accounting for the use of ICA were discussed in section 7.2. However, an unreliability will be present as the ICA algorithm is limited to M = N. The true N is unknown, thus ICA do not guarantee to find all the active sources. Furthermore, the comparison of the main algorithm to ICA was found to be compromised. The localisation and phase of the active sources are not necessary the same for the two estimates, which distorted the MSE between the two matrices to an unknown degree. In despite of this issue, the comparison was still performed by MSE, which suggest a potential to improve the found performance. Against the prior expectation, the an acceptable performance found for the case M = N, though for the cases of M < N the performance was decreased significantly.

Due to the possible unreliability of the performance evaluation with respect to ICA, an alternative test was considered – the alpha wave analysis. However, from the analysis no new conclusion was made. The expected behaviour was not observed, with respect to an increased amount of alpha frequency for the test subject having closed eyes. The behaviour was more or less fluctuating over time. However, exceptions from the expected behaviour were also found for the raw measurement. This indicates that different approaches, with respect to measuring the power within the alpha band, should be considered. The advantage of this test approach, in general, is that one see past the challenge of recovering the exact source signal, but rather consider the practical usage of the source separation. For instance, when considering the usage of the source signals within the a hearing aid, cf. section 1.1.1, the amount of active source signals might be more significant that an exact recovery.

The last issue addressed in this thesis was estimation of the number of active sources k relative to the maximum number of sources N. Through out the implementation of the main algorithm in chapter 6 it was argued that setting k = N would not compromise the results. In fact perhaps lead to better estimates as fewer sources

must be recovered and therefore reduced the errors. This is supported by [4] where the same assumption is used, as no sparsity constraint is considered for Cov-DL. Likewise for M-SBL, providing k would only reduced possible errors within localisation of the sources. However, by setting k = N one must assume that k sources are active, thus no less that k sources are estimated. Hence, to justify this definition of k one must have a qualified guess with respect to the true k. However, this is the issue addressed in the problem statement, as this is not possible in practice.

To address this exact issue an investigation with respect to estimation of k was conducted in chapter 8. Due to interesting empirical observation, it was chosen to analyses the source signal resulting from the main algorithm when $k = N = \overline{M}$. That is estimating the maximum number of active sources under the hypothesis that the false estimates where to be identified among the true estimates. By false estimate there is referred to a non-zero estimate of a zero row. From visual observation of results from the simulated stochastic dataset, a potential was seen as the false estimates appear as copies of the true estimates. Identification of these false estimates was sought by use of the replicates of the true estimates. However, this was not found successful in the desired cases where M < k. Here the false estimates manage to diverge more from the true estimates. Furthermore, the identification method was found to have trouble separating the true estimate from the corresponding replicates. Alternative methods could have been considered with respect to estimation of k. One obvious approach is to consider the estimate of k which could be provided from M-SBL, if a k was not given as a input to the algorithm. However, one could argue that an essential limitation is still present as the mixing matrix is needed for the executing of M-SBL and the number of active sources k is a needed input to the Cov-DL. Furthermore the success rate of such estimation of k was not provided in the article [5] of which the article was based. As the performance presented in the article was obtained by providing k to the algorithm, cf. 5.2.1.

Throughout this chapter the choices which ware found essential toward the obtained conclusions have been discussed and alternative choices have been considered. None of these alternatives are sought investigated in the thesis and but they will serve as an essential point to be considered if further work on the main algorithm where to be conducted.

Chapter 10

Conclusion

The main purpose of this thesis is to prove a state of the art result for solving the EEG inverse problem, with respect to the original brain source signals, in the under-determined case. Secondary the method is sought improved from a practical perspective.

A main algorithm was proposed, based on the state of the art methods covariance-domain dictionary learning (Cov-DL) [4] and multiple sparse Bayesian learning (M-SBL) [5], for recovery of respectively the mixing matrix and the source signals, from the EEG inverse problem. From the initial implementation verification of the Cov-DL method, the method is found to fail. Based on brief analysis of the error, it is concluded that the scientific article proposing the method did not have a sufficient degree of reproducibility. The implementation verification of the M-SBL method is found successful when the true mixing matrix is provided. From this it can be concluded that the corresponding scientific article provided a sufficient degree of reproducibility.

To replace the recovery of the mixing matrix from Cov-DL within, the main algorithm, a fixed matrix was chosen, based on empirical tests. By this modification and the corresponding tests, it is not expected that the main algorithm manages to provide a sufficient recovery of the source signals. The performance of the resulting main algorithm applied to synthetic data is found to be both insufficient and fluctuating, indicating an unreliability which complicates any useful conclusion. Though, with respect to investigating the extent of the resulting main algorithm, the performance was tested on real EEG data and compared to the respective solutions provided by independent component analysis (ICA) on the complete system. From the analysis of the results, a potentially reliable recovery of sources is seen for the complete system, while for the case of interest, the under-determined case, the performance can not be evaluated as sufficient. Thus as expected, it must be concluded that the main algorithm do not provide a sufficient recovery of the source signals. Furthermore, the extent of the algorithm can not be determined based on

the fluctuation results. An alternative test was conducted as an attempt to analysis the recovered source signal from a different perspective. From this analysis no significant founding was discovered, thus the previous conclusion is preserved.

With respect to the practical perspective, the issue of the unknown number of active source relative to the total number of sources was investigate through empirical tests. A method was proposed with respect to identification of non-active sources after the recovery. The method showed potential when applied to results provided by use of the true mixing matrix. However, when increasing the over-completeness as desired difficulties arise. Thus, it is concluded that the method does not provide a sufficient estimate of the number of active sources. But the found potential suggests that further work on the method may allow the possibility of reaching a reliable estimate.

Overall, it is concluded that a recreation of the specified state of the art methods for source recovery was not successfully provided by the proposed main algorithm. Furthermore, the proposed alternative to the estimation of the mixing matrix did not provide reliable results hence the conclusion remains. Secondary it is concluded that a potential is seen with respect to using the M-SBL method in practice, considering the unknown number of active sources.

Chapter 11

Further Studies

Based on the accomplished conclusions, further studies regarding the proposed main algorithm and the general issue of source signal recovery will be discussed.

One essential finding in this thesis was the negative result with respect to the proposed implementation of the COV-DL method. It is concluded that the reproducibility of the corresponding article was not sufficient. However, it is not excluded that further studies would enable a successful implementation. This could be further investigations in form of rewriting the optimisation problem or choosing a different optimisation process.

Another aspect could be to dismiss the COV-DL method and do some research with respect to alternative methods for finding the mixing matrix. Such method could be the low resolution electrical tomograph (LORETA), which localises electrical activity inside the brain. Or, the minimum norm estimates (MNE), which reconstructs the activity on the cortical surface [??].

From the overall perspective of the topic, it could be of interest to alter the view on the EEG measurements. In this thesis the purpose was to recover active source signals from the EEG measurements, by the main algorithm. A news article from April 2020, [18], presents the newest research from Eriksholm Research Center. Concerning application of EEG measurements within a hearing aid, as mentioned in the chapter 1. They suggest that, by focusing only on recovery of signals from eye movements, the direction of the sight can be measured. As this thesis focus on finding all sources from the provided EEG measurements, this could be a change of focus. The advantage of targeting the specific signals, which was before considered as noise on the EEG measurement, is that fewer sensors is needed and fewer signals are sought recovered. This results in fewer computations and a potential of avoiding the difficult under-determined case could be present. For this case a different EEG measurements data base must be provided or created since the data used in this thesis do not contain the signals created by the eye movement and surrounding muscle movements.

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Appendix A

Supplementary Theory for Chapter 4

Throughout this chapter supplementary theory for understanding the covariance-domain dictionary learning (Cov-DL) is described. First an introduction to compressive sensing which is the framework behind Cov-DL. Then an dictionary learning algorithm used for finding the dictionary matrix \mathbf{D} in section 4.2.1 will be described. And at last principal component analysis (PCA) is introduced as the method behind finding \mathbf{D} in section 4.2.2.

Læs denne introduktio

A.1 Introduction to Compressive Sensing

Compressive sensing is the theory of efficient recovery of a signal from a minimal number of observed measurements. It is build upon empirical observations assuring that many signals can be approximated by remarkably sparser signals. Assume linear acquisition of the observed measurements, then the relation between the measurements and the signal to be recovered can be modelled by the multiple measurement vector (MMV) model (3.2) [15].

Through this section the introduction of the theory behind compressive sensing will be presented for one measurement vector of (3.2), \mathbf{y} , such that the theory is based on the linear system (3.1). This will be done for simplicity, but the theory will still apply for the extended linear system (3.2).

In compressive sensing terminology, $\mathbf{x} \in \mathbb{R}^N$ is the signal of interest sought recovered from the EEG measurement $\mathbf{y} \in \mathbb{R}^M$ by solving the linear system (3.1). In the typical compressive sensing case, the system is under-determined, M < N, and there will therefore exist infinitely many solutions, provided that one solution exist. However, by enforcing certain sparsity constraints it is possible to recover the wanted signal, hence the term sparse signal recovery [15]. The sparsity constraints are the ones presented in 3.1 where the ℓ_0 is introduced to count the non-zeros of the signal

of interest, the source vector \mathbf{x} . The number of non-zeros (active sources) k describe how sparse the source vector is.

To find a k-sparse solution to the linear system (3.1) it can be viewed as the following optimisation problem.

$$\mathbf{x}^* = \arg\min_{\mathbf{x} \in C} \|\mathbf{x}\|_0$$
 subject to $\mathbf{A}\mathbf{x} = \mathbf{y}$.

Unfortunately, this optimisation problem is non-convex due to the definition of the ℓ_0 -norm and is therefore difficult to solve – it is a NP-hard problem. Instead, by replacing the ℓ_0 -norm with the ℓ_1 -norm, the optimisation problem can be approximated and hence becomes computationally feasible [10, p. 27]

$$\mathbf{x}^* = \arg\min_{\mathbf{x} \in C} \|\mathbf{x}\|_1$$
 subject to $\mathbf{A}\mathbf{x} = \mathbf{y}$. (A.1)

With this optimisation problem the best k-sparse solution \mathbf{x}^* can be found. The optimisation problem is referred to as ℓ_1 optimisation problem or Basis Pursuit. The following theorem justifies that the ℓ_1 optimisation problem finds a sparse solution [15, p. 62-63].

A mixing matrix $\mathbf{A} \in \mathbb{R}^{M \times N}$ is defined with columns $\mathbf{A} = [\mathbf{a}_1, \dots, \mathbf{a}_N]$. By assuming uniqueness of a solution \mathbf{x}^* to

$$\min_{\mathbf{x} \in \mathbb{R}^N} \|\mathbf{x}\|_1 \quad \text{subject to} \quad \mathbf{A}\mathbf{x} = \mathbf{y},$$

 $\begin{aligned} \min_{\mathbf{x} \in \mathbb{R}^N} \|\mathbf{x}\|_1 \quad \text{subject to} \quad \mathbf{A}\mathbf{x} = \mathbf{y}, \\ \text{the system } \{\mathbf{a}_j, j \in \text{supp}(\mathbf{x}^*)\} \text{ is linearly independent, and in particular} \\ \|\mathbf{x}^*\|_0 = \operatorname{card}(\operatorname{supp}(\mathbf{x}^*)) \leq M. \end{aligned}$

$$\|\mathbf{x}^*\|_0 = \operatorname{card}(\operatorname{supp}(\mathbf{x}^*)) \leq M$$

Proof

Assume that the set $\{\mathbf{a}_l, l \in S\}$ of l columns from matrix $\mathbf{A} \in \mathbb{R}^{M \times N}$ is linearly dependent with the support $S = \text{supp}(\mathbf{x}^*)$. Thus a non-zero vector $\mathbf{v} \in \mathbb{R}^N$ supported on S exists such that $\mathbf{A}\mathbf{v} = \mathbf{0}$ – the system is linear dependent. The unique solution \mathbf{x}^* can then be written as, for any $t \neq 0$,

$$\|\mathbf{x}^*\|_1 < \|\mathbf{x}^* + t\mathbf{v}\|_1 = \sum_{l \in S} |x_l^* + tv_l| = \sum_{l \in S} \operatorname{sgn}(x_l^* + tv_l)(x_l^* + tv_l). \tag{A.2}$$

For a small |t|

$$|t| < \min_{l \in S} \frac{|x_l^*|}{\|\mathbf{v}\|_{\infty}},$$

then the sign function becomes

$$\operatorname{sgn}(x_l^* + tv_l) = \operatorname{sgn}(x_l^*), \quad \forall l \in S.$$

By including this result in (A.2) and remembering $t \neq 0$:

$$\|\mathbf{x}^*\|_1 < \sum_{l \in S} \operatorname{sgn}(x_l^*)(x_l^* + tv_l) = \sum_{l \in S} \operatorname{sgn}(x_l^*)x_l^* + t\sum_{l \in S} \operatorname{sgn}(x_l^*)v_l = \|\mathbf{x}^*\|_1 + t\sum_{l \in S} \operatorname{sgn}(x_l^*)v_l.$$

From this it can be seen that it is always possible to choose $t \neq 0$ small enough such that

$$t\sum_{l\in S}\operatorname{sgn}(x_l^*)v_l\leq 0,$$

which contradicts that \mathbf{v} make the columns of \mathbf{A} linear dependent. Therefore, the set $\{\mathbf{a}_l, l \in S\}$ must be linearly independent.

From the theorem it must be concluded that the choice of the mixing matrix \mathbf{A} has a significant impact on whenever a unique solution \mathbf{x}^* exist for the ℓ_1 optimisation problem (A.1). Therefore, when recovering \mathbf{A} , some considerations regarding the recovering process of \mathbf{A} must be taken into account. A method for the recovering of \mathbf{A} could be to use a dictionary. This will be explained in the following section 4.2.1.

An alternative solution method to the ℓ_1 optimisation includes greedy algorithms such as the Orthogonal Matching Pursuit (OMP) [15, P. 65]. The OMP algorithm is an iteration process where an index set S is updated – at each iteration – by adding indices corresponding to the columns of \mathbf{A} which describe the residual best possible, hence greedy. The vector \mathbf{x} is then updated by a vector supported on S which minimise the residual, that is the orthogonal projection of \mathbf{y} onto the span $\{\mathbf{a}_l \mid l \in S\}$.

A.2 K-SVD Algorithm

The dictionary learning algorithm K-SVD provides an updating rule which is applied to each column of $\mathbf{A}_0 = [\mathbf{a}_1, \dots, \mathbf{a}_N]$ where \mathbf{A}_0 being a random initial dictionary matrix. Updating first \mathbf{a}_j for $j = 1, \dots, N$ and then the corresponding row of \mathbf{X} , \mathbf{x}_i . for j = i. Let \mathbf{a}_{j_0} be the column to be updated and let the remaining columns be fixed. By rewriting the objective function in (4.5) using matrix notation it is possible to isolate the contribution from \mathbf{a}_{j_0} .

$$\|\mathbf{Y} - \mathbf{A}\mathbf{X}\|_{F}^{2} = \|\mathbf{Y} - \sum_{\substack{j=1\\j=i}}^{N} \mathbf{a}_{j} \mathbf{x}_{i} \|_{F}^{2}$$

$$= \|\left(\mathbf{Y} - \sum_{\substack{j=j\\j\neq i\\j=i}}^{N} \mathbf{a}_{j} \mathbf{x}_{i} - \mathbf{a}_{j_{0}} \mathbf{x}_{i_{0}} \right) - \mathbf{a}_{j_{0}} \mathbf{x}_{i_{0}} \|_{F}^{2}, \tag{A.3}$$

where i = j, $i_0 = j_0$ and where F is the Frobenius norm that works on matrices

$$\|\mathbf{A}\|_F = \sqrt{\sum_{i=1}^M \sum_{j=1}^N |a_{ij}|^2}.$$

In (A.3) the term in the parenthesis is denoted by \mathbf{E}_{j_0} , an error matrix, and hence by minimising (A.3) with respect to \mathbf{a}_{j_0} and \mathbf{x}_{i_0} leads to the optimal contribution from j_0

$$\min_{\mathbf{a}_{j_0}, \mathbf{x}_{i_0}.} \| \mathbf{E}_{j_0} - \mathbf{a}_{j_0} \mathbf{x}_{i_0}. \|_F^2.$$
 (A.4)

The optimal solution to (A.4) is known to be the rank-1 approximation of \mathbf{E}_{j_0} [14, p. 232]. That is a partial single value decomposition (SVD) makes the best low-rank approximation of a matrix such as \mathbf{E}_{j_0} . The SVD is given as

$$\mathbf{E}_{j_0} = \mathbf{U} \mathbf{\Sigma} \mathbf{V}^T \in \mathbb{R}^{M \times N},$$

with $\mathbf{U} \in \mathbb{R}^{M \times M}$ and $\mathbf{V} \in \mathbb{R}^{N \times N}$ being unitary matrices¹ and $\mathbf{\Sigma} = \operatorname{diag} [\boldsymbol{\sigma}_1, \dots, \boldsymbol{\sigma}_M] \in \mathbb{R}^{M \times N}$ a diagonal matrix. $\boldsymbol{\sigma}_j$ are the non-negative singular values of \mathbf{E}_{j_0} . The best k-rank approximation to \mathbf{E}_{j_0} , with $k < \operatorname{rank}(\mathbf{E}_{j_0})$ is then given by [14, p. 232]:

$$\mathbf{E}_{j_0}^{(k)} = \sum_{j=1}^k \boldsymbol{\sigma}_j \mathbf{u}_j \mathbf{v}_j^T.$$

Since the outer product always has rank-1 letting $\mathbf{a}_{j_0} = \mathbf{u}_1$ and $\mathbf{x}_{i_0} = \boldsymbol{\sigma}_j \mathbf{v}_1^T$ solves the optimisation problem (A.4). However in order to preserve the sparsity in \mathbf{X} while optimising, only the non-zero entries in \mathbf{x}_{i_0} are allowed to vary. For this purpose only a subset of columns in \mathbf{E}_{j_0} is considered, those which correspond to the non-zero entries of \mathbf{x}_{i_0} . A matrix \mathbf{P}_{i_0} is defined to restrict \mathbf{x}_{i_0} to only contain the non-zero-rows corresponding to N_{j_0} non-zero rows:

$$\mathbf{x}_{i_0}^{(R)} = \mathbf{x}_{i_0}.\mathbf{P}_{i_0}$$

where R denoted the restriction. By applying the SVD to the error matrix which has been restricted $\mathbf{E}_{j_0}^{(R)} = \mathbf{E}_{j_0} \mathbf{P}_{i_0}$ and updating \mathbf{a}_{j_0} and $\mathbf{x}_{i_0}^{(R)}$ the rank-1 approximation is found and the original representation vector is updated as $\mathbf{x}_{i_0} = \mathbf{x}_{i_0}^{(R)} \mathbf{P}_{i_0}^T$.

The main steps of K-SVD is described in algorithm 3.

¹Unitary matrix: $\mathbf{U}^T\mathbf{U} = \mathbf{U}\mathbf{U}^T = \mathbf{I}$

Algorithm 3 K-SVD

```
1: k = 0
  2: Initialize random A_{(0)}
  3: Initialize \mathbf{X}_{(0)} = \mathbf{0}
       procedure K\text{-SVD}(\mathbf{A}_{(0)})
               Normalize columns of A_{(0)}
  6:
               while error \geq limit do
  7:
                      j = j + 1
  8:
                      for j \leftarrow 1, 2, \dots, L do
  9:
                                                                                                                     \triangleright updating each col. in \mathbf{X}_{(k)}
                             \hat{\mathbf{x}}_j = \min_{\mathbf{x}} \|\mathbf{y}_j - \mathbf{A}_{(k-1)}\mathbf{x}_j\| subject to \|\mathbf{x}_j\| \le k > use Basis Pursuit
10:
11:
                      end for
12:
                      \mathbf{X}_{(k)} = {\{\hat{\mathbf{x}}_j\}_{j=1}^L}
                      for j_0 \leftarrow 1, 2, \cdots, N do
13:
                              \Omega_{j_0} = \{ j \mid 1 \le j \le L, \mathbf{X}_{(k)}[j_0, j] \ne 0 \}
14:
                              From \Omega_{i_0} define \mathbf{P}_{i_0}
15:
                             \mathbf{E}_{j_0} = \mathbf{Y} - \sum_{j \neq j_0}^{N} \mathbf{a}_j \mathbf{x}_i.
16:
                             \mathbf{E}_{j_0}^{(R)} = \mathbf{E}_{j_0} \mathbf{P}_{i_0}\mathbf{E}_{j_0}^{(R)} = \mathbf{U} \mathbf{\Sigma} \mathbf{V}^T
17:
                                                                                                                                                  ▶ perform SVD
18:
                              \mathbf{a}_{j_0} \leftarrow \mathbf{u}_1
                                                                                                                       \triangleright update the j_0 col. in \mathbf{A}_{(k)}
19:
                              (\mathbf{x}_{i_0})^{(R)} \leftarrow \boldsymbol{\sigma}_1 \mathbf{v}_1 
 \mathbf{x}_{i_0} \leftarrow (\mathbf{x}_{i_0})^{(R)} \mathbf{P}_{i_0}^T 
20:
                                                                                                                       \triangleright update the i_0 row in \mathbf{X}_{(k)}
21:
                       end for
22:
                      error = \|\mathbf{Y} - \mathbf{A}_{(k)} \mathbf{X}_{(k)}\|_F^2
23:
               end while
25: end procedure
```

A.3 Principal Component Analysis

In this section the method behind principal component analysis (PCA) used for the Cov-DL described in section 4.2.2.

PCA is dimensionality reduction method used for reduction of dimension of large datasets. In short, PCA used the statistical information of mean, variance and correlation between the data to transform the large dataset into smaller datasets while maintaining most of the original information. These smaller datasets are known as the principal components and contain most of the information of the dataset but with fewer dimensions. For some datasets, before PCA is applied, the data must undergo a standardization/scaling to remove any difference in the data. This is essential as large differences between the data would dominate. The standardization

of a dataset **Z** is performed by

$$\tilde{\mathbf{z}}_i = \frac{\mathbf{z}_i - \bar{\mathbf{z}}_i}{s_{\mathbf{z}_i}}, \quad \forall i = 1, \dots, m$$

where \mathbf{z}_i is a row of a matrix \mathbf{Z} , $\bar{\mathbf{z}}_i$ is the sample mean of \mathbf{z}_i and $s_{\mathbf{z}_i}$ is the standard deviation of \mathbf{z}_i . The standardized dataset is now giving by $\tilde{\mathbf{Z}}$. The standardization step is not necessary in the case of real EEG scalp measurements as no large difference between the data is present.

With \mathbf{Z} or the scaled data $\tilde{\mathbf{Z}}$ a correlation matrix is computed as

$$\Sigma_{\mathbf{Z}} = \operatorname{Corr}(\mathbf{Z}) = \frac{1}{m} \mathbf{Z}^T \mathbf{Z}.$$

From the correlation matrix a orthonormal basis of eigenvectors $\mathbf{p}_1, \dots, \mathbf{p}_m$ with corresponding eigenvalues $\lambda_1, \dots, \lambda_m$ exists, cf. theorem 6.15 in [21, p. 375]. Furthermore, one assume that $\lambda_1 \geq \dots \geq \lambda_m$. The eigenvectors and eigenvalues can be computed from the correlation matrix by e.g. using a singular value decomposition (SVD). With SVD an orthogonal matrix \mathbf{P} with the eigenvectors $\mathbf{p}_1, \dots, \mathbf{p}_m$ as columns is obtain with the associated eigenvalues as a diagonal matrix denoted as \mathbf{P}_{diag} . The principal components is then defined by

$$\mathbf{u}_i = \mathbf{Z}\mathbf{p}_i$$

where \mathbf{p}_i is the *i*-th eigenvector of the correlation matrix $\Sigma_{\mathbf{Z}}$. Thus each principal component is a linear combination of the dataset \mathbf{Z} [21, p. 460]. With the principal components the first N components forms a set of basis vectors $\mathbf{U} = [\mathbf{u}_1, \dots, \mathbf{u}_N]$.

Appendix B

Derivations for Multiple Sparse Bayesian Learning

B.1 Derivation of Posterior Mean \mathcal{M} and Covariance Σ

The purpose of this section is to derive the mean \mathcal{M} and covariance Σ of the posterior distribution

$$p\left(\mathbf{x}_{\cdot j}|\mathbf{y}_{\cdot j};\boldsymbol{\gamma}\right) \sim \mathcal{N}(\boldsymbol{\mu}_{\cdot j},\boldsymbol{\Sigma}).$$

from (5.2) in section 5.1.1.

Let $\mathbf{x}_{\cdot j} = \mathbf{x}$ and $\mathbf{y}_{\cdot j} = \mathbf{y}$ to ease the notation throughout the derivation. The prior and likelihood is then defined as

$$p(\mathbf{x}; \boldsymbol{\gamma}) \sim \mathcal{N}(\mathbf{0}, \boldsymbol{\gamma} \mathbf{I})$$

$$p(\mathbf{y}|\mathbf{x}) \sim \mathcal{N}(\mathbf{A}\mathbf{x}, \sigma^2 \mathbf{I})$$

Now define one Gaussian random variable

$$\mathbf{z} = \begin{bmatrix} \mathbf{x} \\ \mathbf{y} | \mathbf{x} \end{bmatrix} \in \mathbb{R}^{N+M}.$$

The mean and covariance of the Gaussian variable ${\bf z}$ can then be partitioned into

$$\Sigma_{\mathbf{z}} = \begin{bmatrix} \Sigma_{\mathbf{x}\mathbf{x}} & \Sigma_{\mathbf{x}\mathbf{y}} \\ \Sigma_{\mathbf{y}\mathbf{x}} & \Sigma_{\mathbf{y}\mathbf{y}} \end{bmatrix} \in \mathbb{R}^{N+M \times N+M}$$

and

$$\mu_{\mathbf{z}} = \begin{bmatrix} \mu_{\mathbf{x}} \\ \mu_{\mathbf{y}|\mathbf{x}} \end{bmatrix} = \begin{bmatrix} 0 \\ \mathbf{A}\mathbf{x} \end{bmatrix}.$$

Then, from [13], the posterior distribution of $p(\mathbf{x}_{\cdot j}|\mathbf{y}_{\cdot j}; \boldsymbol{\gamma})$ is defined but the covariance for \mathbf{y} is not known. Instead, one assume that the covariance for \mathbf{y} given \mathbf{x} can be used instead:

$$\Sigma = cov(\mathbf{x}, \mathbf{x}|\mathbf{y}) = \Sigma_{\mathbf{x}\mathbf{x}} - \Sigma_{\mathbf{x}\mathbf{y}} \Sigma_{\mathbf{y}\mathbf{y}|\mathbf{x}}^{-1} \Sigma_{\mathbf{y}\mathbf{x}}$$
$$\mu = \mu_{\mathbf{x}} + \Sigma_{\mathbf{x}\mathbf{y}} \Sigma_{\mathbf{y}\mathbf{y}|\mathbf{x}}^{-1} (\mathbf{y} - \mu_{\mathbf{y}})$$

Each of the conditional covariances within the previous expressing will now be found.

Conditional Covariance Σ_{xx} The covariance of x comes directly from the distribution

$$\Sigma_{xx} = \gamma I$$

Conditional Covariance Σ_{yx} The covariance between x and y|x is found by using the linearity of covariance and y = Ax + e:

$$\begin{split} \boldsymbol{\Sigma}_{\mathbf{y}\mathbf{x}} &= \operatorname{cov}\left(\mathbf{x}, \mathbf{A}\mathbf{x} + \mathbf{e}\right) \\ &= \operatorname{cov}\left(\mathbf{x}, \mathbf{A}\mathbf{x}\right) + \operatorname{cov}\left(\mathbf{x}, \mathbf{e}\right) \\ &= \boldsymbol{\Sigma}_{\mathbf{x}\mathbf{x}} \mathbf{A}^T \\ &= \boldsymbol{\gamma} \mathbf{I} \mathbf{A}^T \end{split}$$

where $cov(\mathbf{x}, \mathbf{e}) = 0$ because they are uncorrelated.

Conditional Covariance $\Sigma_{yy|x}$ Lastly the covariance of y|x is similarly found by the definition of conditional covariance under the assumption that $\Sigma_{yy} = \sigma^2 I$:

$$\Sigma_{\mathbf{y}\mathbf{y}|\mathbf{x}} = \operatorname{cov}(\mathbf{y}, \mathbf{y}|\mathbf{x}) = \Sigma_{\mathbf{y}\mathbf{y}} - \Sigma_{\mathbf{y}\mathbf{x}} \Sigma_{\mathbf{x}\mathbf{x}}^{-1} \Sigma_{\mathbf{x}\mathbf{y}}$$
$$= \sigma^{2} \mathbf{I} - \gamma \mathbf{I} \mathbf{A}^{T} (\gamma \mathbf{I})^{-1} \mathbf{A} \gamma \mathbf{I}$$
$$= \sigma^{2} \mathbf{I} - \mathbf{A} \gamma \mathbf{I} \mathbf{A}^{T}.$$

By combining all the found conditional covariances the resulting covariance becomes

$$\boldsymbol{\Sigma} = \boldsymbol{\gamma} \mathbf{I} - \boldsymbol{\gamma} \mathbf{I} \mathbf{A}^T (\boldsymbol{\Sigma}_{\mathbf{y}\mathbf{y}|\mathbf{x}})^{-1} \mathbf{A} \boldsymbol{\gamma} \mathbf{I}.$$

The resulting mean under the assumption that $\mu_{y} = 0$ becomes

$$\mu_{\cdot,j} = \mathbf{0} + \gamma \mathbf{I} \mathbf{A}^T (\mathbf{\Sigma}_{yy})^{-1} (\mathbf{y} - \mathbf{0})$$
$$= \gamma \mathbf{I} \mathbf{A}^T (\mathbf{\Sigma}_{yy})^{-1} \mathbf{y}$$

Appendix C

Independent Component Analysis

This appendix provide the basic theory of independent component analysis (ICA). The theory is necessary if one wants a deeper understanding towards the justification of applying result from ICA as a reference for evaluation of the main algorithm proposed in this thesis. The appendix concludes with an algorithm specifying ICA method which is applied in the thesis. Additionally, an verification test is conducted to evaluate the applied ICA method on the synthetic data, cf. 6.2.

C.1 Basic Theory of Independent Component Analysis

Independent component analysis (ICA) is a method that applies to the general problem of decomposition of a measurement vector into a source vector and a mixing matrix. The intention of ICA is to separate a multivariate signal into statistical independent and non-Gaussian signals, and identify the mixing matrix \mathbf{A} , given only the observed measurements \mathbf{Y} . A well known application example of source separation is the cocktail party problem, where it is sought to listen to one specific person speaking in a room full of people having interfering conversations. Let $\mathbf{y} \in \mathbb{R}^M$ be a single measurement from M microphones containing a linear mixture of all the speak signals that are present in the room. When additional noise is not considered, the problem can be described as the familiar linear system

$$y = Ax \tag{C.1}$$

where $\mathbf{x} \in \mathbb{R}^N$ contain the N underlying speak signals and \mathbf{A} is a mixing matrix where the coefficients may depend on the distances from a source to the microphone. As such each y_i is a weighted sum of all the sources of speak present to the i-th microphone. By ICA both the mixing matrix \mathbf{A} and the source signals \mathbf{x} are sought estimated from the observed measurements \mathbf{y} . The main attribute of ICA is the

assumption that the sources in \mathbf{x} are statistically independent and non-Gaussian distributed, hence the name independent components.

By independence, one means in general that changes in one source signal do not affect the other source signals. In theory n variables x_1, \dots, x_n is independent if the joint probability density function (pdf) of \mathbf{x} satisfies

$$p(x_1, x_2, ..., x_n) = p_1(x_1)p_2(x_2)\cdots p_n(x_n).$$

The possibility of separating a signal into independent and non-Gaussian components originates from the central limit theorem [19, p. 34]. The theorem states that the distribution of any linear mixture of two or more independent random variables tends toward a Gaussian distribution, under certain conditions. For instance the distribution of a mixture of two independent random variables is always closer to a Gaussian distribution than the original variables. In other word the original variables is most non-Gaussian. The application of the central limit theorem within ICA will be elaborated later in this appendix.

C.1.1 Assumptions and Preprocessing

For simplicity assume **A** is square i.e. M = N and invertible. As such when **A** has been estimated the inverse is computed and the components can simply be estimated as $\mathbf{x} = \mathbf{A}^{-1}\mathbf{y}$ [19, p. 152-153]. As both **A** and **x** are unknown the variances of the independent components can not be determined. However it is reasonable to assume that **x** has unit variance – **A** is assume to have unit variance as well. Any scalar multiplier within a source can be cancelled out by dividing the corresponding column in **A** with the same scalar [19, p. 154]. For further simplification it is assumed without loss of generality that $\mathbb{E}[\mathbf{y}] = 0$ and $\mathbb{E}[\mathbf{x}] = 0[19, p. 154]$. In case this assumption is not true, the measurements can be centred by subtracting the mean as preprocessing before doing ICA.

A preprocessing step central to ICA is to whiten the measurements \mathbf{y} . By the whitening process any correlation in the measurements are removed and unit variance is ensured – the independent components \mathbf{x} becomes uncorrelated and have unit variance. Furthermore, this reduces the complexity of ICA and therefore simplifies the recovering process. Whitening is a linear transformation of the observed data. This is multiplying the measurement vector \mathbf{y} with a whitening matrix \mathbf{V} ,

$$\mathbf{y}_{\mathrm{white}} = \mathbf{V}\mathbf{y}$$

to obtain a new measurement vector \mathbf{y}_{white} which is considered white. To obtain a whitening matrix the eigenvalue decomposition (EVD) of the covariance matrix can be used,

$$\mathbb{E}[\mathbf{y}\mathbf{y}^T] = \mathbf{E}\mathbf{D}\mathbf{E}^T$$

where \mathbf{D} is a diagonal matrix of eigenvalues and \mathbf{E} is a matrix consisting of the associated eigenvectors. From \mathbf{E} and \mathbf{D} a whitening matrix \mathbf{V} is constructed as

$$\mathbf{V} = \mathbf{E}\mathbf{D}^{-1/2}\mathbf{E}^T.$$

where $\mathbf{D}^{-1/2} = \operatorname{diag}(d_1^{-1/2}, \dots, d_n^{-1/2})$ is a component-wise operation[19, p.159]. By multiplying the measurement vector \mathbf{y} with a whitening matrix \mathbf{V} the data becomes white

$$\mathbf{y}_{\text{white}} = \mathbf{V}\mathbf{y} = \mathbf{V}\mathbf{A}\mathbf{x} = \mathbf{A}_{\text{white}}\mathbf{x}.$$

Furthermore the corresponding mixing matrix $\mathbf{A}_{\text{white}}$ becomes orthogonal

$$\mathbb{E}[\mathbf{y}_{\text{white}}\mathbf{y}_{\text{white}}^T] = \mathbf{A}_{\text{white}}\mathbb{E}[\mathbf{x}\mathbf{x}^T]\mathbf{A}_{\text{white}}^T = \mathbf{A}_{\text{white}}\mathbf{A}_{\text{white}}^T = \mathbf{I},$$

where $\mathbb{E}[\mathbf{x}\mathbf{x}^T] = \mathbf{I}$ due to \mathbf{x} having zero mean and uncorrelated entries.

As a consequence ICA can restrict its search for the mixing matrix to the orthogonal matrix space – that is instead of estimating N^2 parameters ICA has only to estimate an orthogonal matrix which has N(N-1)/2 parameters [19, p. 159].

C.1.2 Recovery of the Independent Components

The estimation of the mixing coefficients a_{ij} and the independent components x_i by ICA is now elaborated, based on [19, p. 166].

The simple and intuitive method is to take advantage of the assumption of non-Gaussian independent components. Consider again the model of a single measurement vector $\mathbf{y} = \mathbf{A}\mathbf{x}$, where the data vector comply to the assumption of being mixture of independent components. Here the independent components can be estimated by the inverted model

$$\mathbf{x} = \mathbf{A}^{-1}\mathbf{y}.$$

Consider first the estimation of a single independent component x_i , here a linear combination of y_i is considered. Denote for now a single independent component by z such that

$$z = \mathbf{b}^T \mathbf{y} = \sum_k b_k y_k \tag{C.2}$$

where one want to determine the vector \mathbf{b} . This can be rewritten to

$$z = \mathbf{b}^T \mathbf{A} \mathbf{x}$$

From this it is seen that y is a linear combination of the x_i with coefficients given by the vector $\mathbf{b}^T \mathbf{A}$, now denote this vector by \mathbf{q} . As such

$$z = \mathbf{b}^T \mathbf{y} = \mathbf{b}^T \mathbf{A} \mathbf{x} = \mathbf{q}^T \mathbf{x} = \sum_k q_k x_k.$$
 (C.3)

By this expression, consider the thought of **b** being one of the rows in \mathbf{A}^{-1} , then the linear combination $\mathbf{b}^T \mathbf{y}$ is equal to one of the independent components.

The objective is now to apply the central limit theorem to determine \mathbf{b} such that it equals one of the rows of \mathbf{A}^{-1} . As \mathbf{A} is unknown it is not possible to determine \mathbf{b} exactly, but an estimate can be found to make a good approximation.

Due to z denoting some x_i it is clear that the equality in (C.3) only holds true when \mathbf{q} consist of only one non-zero element equal to 1. Thus, from the central limit theorem the distribution of $\mathbf{q}^T\mathbf{x}$ is most non-Gaussian when it equals one of the independent components which was assumed non-Gaussian. As such, since $\mathbf{q}^T\mathbf{x} = \mathbf{b}^T\mathbf{y}$, it is possible to vary the coefficients in \mathbf{b} and look at the distribution of $\mathbf{b}_i.\mathbf{y}$. Finding the vector \mathbf{b}^T that maximizes the non-Gaussianity would then correspond to $\mathbf{q} = \mathbf{A}^T\mathbf{b}^T$ having only a single non-zero element. Thus maximizing the non-Gaussianity of $\mathbf{b}_i.\mathbf{y}$ results in one of the independent components [19, p. 166]. Considering the N-dimensional space of vectors \mathbf{b}^T there exist 2N local maxima, corresponding to x_i and $-x_i$ for all N independent components [19, p. 166].

C.1.3 Kurtosis

To maximize the non-Gaussianity a measure for Gaussianity is needed. Kurtosis is a quantitative measure used for non-Gaussianity of random variables [19, p.171]. Kurtosis of a random variable y is the fourth-order cumulant denoted by kurt(y). For y with zero mean and unit variance, kurtosis reduces to

$$\operatorname{kurt}(y) = \mathbb{E}[y^4] - 3.$$

It is seen that the kurtosis is a normalized version of the fourth-order moment defined as $\mathbb{E}[y^4]$. For a Gaussian random variable the fourth-order moment equals $3(\mathbb{E}[y^2])^2$ hence the corresponding kurtosis will be zero [19, p. 171]. Consequently the kurtosis of non-Gaussian random variables will almost always be different from zero.

The kurtosis is a common measure for non-Gaussianity due to its simplicity both theoretical and computational. The kurtosis can be estimated computationally by the fourth-order moment of sample data when the variance is constant. Furthermore, for two independent random variables x_1, x_2 the following linear properties applies to the kurtosis of the sum

$$\operatorname{kurt}(x_1 + x_2) = \operatorname{kurt}(x_1) + \operatorname{kurt}(x_2)$$
 and $\operatorname{kurt}(\alpha x_1) = \alpha^4 \operatorname{kurt}(x_1)$

However, one complication concerning kurtosis as a measure is that kurtosis is sensitive to outliers [19, p. 182].

Consider from (C.3) the vector $\mathbf{q} = \mathbf{A}^T \mathbf{b}$ such that $\mathbf{b}^T \mathbf{y} = \sum_{k=1}^{\infty} q_k x_k$. By the additive property of kurtosis

$$\operatorname{kurt}\left(\mathbf{b}^{T}\mathbf{y}\right) = \sum_{k=1}^{\infty} q_{k}^{4} \operatorname{kurt}(x_{k}).$$

Then the assumption of the independent components having unit variance results in $\mathbb{E}[x_i^2] = \sum_{k=1} q_k^2 = 1$. That is geometrically that **q** is constrained to the unit sphere, $\|\mathbf{q}\|^2 = 1$.

By this an optimisation problem maximising the kurtosis of $\mathbf{b}^T \mathbf{y}$ is similar to maximizing $|\text{kurt}(x_i)| = |\sum_{k=1} q_k^4 \text{kurt}(x_k)|$ on the unit sphere. Due to the described preprocessing \mathbf{b}^T is assumed to be white and it can be shown that $\|\mathbf{q}\| = \|\mathbf{b}^T\|$ [19, p. 174]. This shows that constraining $\|\mathbf{q}\|$ to one is similar to constraining $\|\mathbf{b}^T\|$ to one.

C.1.4 Basic ICA algorithm

Now a basic ICA algorithm is specified, this algorithm is based on the gradient optimisation method with kurtosis.

The general idea behind a gradient algorithm is to determine the direction for which $\operatorname{kurt}(\mathbf{b}^T\mathbf{y})$ is growing the most, based on the gradient. The gradient of $|\operatorname{kurt}(\mathbf{b}^T\mathbf{y})|$ is computed as

$$\frac{\partial |\text{kurt}(\mathbf{b}^T \mathbf{y})|}{\partial \mathbf{b}} = 4 \text{sign}(\text{kurt}(\mathbf{b}^T \mathbf{y})) (\mathbb{E}[\mathbf{y}(\mathbf{b}^T \mathbf{y})^3] - 3\mathbf{y}\mathbb{E}[(\mathbf{b}^T \mathbf{y})^2])$$
(C.4)

As $\mathbb{E}[(\mathbf{b}^T \mathbf{y})^2] = \|\mathbf{y}\|^2$ for whitened data the corresponding term does only affect the norm of \mathbf{b} within the gradient algorithm. Thus, as it is only the direction that is of interest, this term can be omitted. Because the optimisation is restricted to the unit sphere a projection of \mathbf{b} onto the unit sphere must be performed in every step of the gradient method. This is done by dividing \mathbf{b} by its norm. This gives update step [19, p. 178]

$$\Delta \mathbf{b} \propto \operatorname{sign} \left(\operatorname{kurt}(\mathbf{b}^T \mathbf{y}) \right) \mathbb{E} [\mathbf{y} (\mathbf{b}^T \mathbf{y})^3]$$
$$\mathbf{b} \leftarrow \mathbf{b} / \|\mathbf{b}\|$$

The expectation operator can be omitted in order to achieve an adaptive version of the algorithm, now using every measurement \mathbf{y} . However, the expectation operator from the definition of kurtosis can not be omitted and must therefore be estimated. This can be done by γ by serving it as the learning rate of the gradient method.

$$\Delta \gamma \propto ((\mathbf{b}^T \mathbf{y})^4 - 3) - \gamma$$

Algorithm 4 combines the above theory, to give an overview of the basic ICA procedure.

Algorithm 4 Basis ICA

```
1: procedure Pre-processing(\mathbf{y})
           Center measurements \mathbf{y} \leftarrow \mathbf{y} - \bar{\mathbf{y}}
 2:
           Whitening \mathbf{y} \leftarrow \mathbf{y}_{white}
 3:
     end procedure
 5:
     procedure ICA(y)
 6:
           k = 0
 7:
 8:
           Initialise random vector
                                                                                                                       ▶ unit norm
           Initialise random value \gamma_{(k)}
 9:
           for j \leftarrow 1, 2, \dots, N do
10:
                 while convergence critia not meet do
11:
                       k = k + 1
12:
                       \mathbf{b}_{(k)} \leftarrow \operatorname{sign} \gamma_{(k-1)} \mathbf{y} (\mathbf{b}_{(k-1)} \mathbf{y})^3
13:
                      \mathbf{b}_{(k)} \leftarrow \mathbf{b}_{(k)} / \|\mathbf{b}_{(k)}\|
14:
                      \gamma_{(k)} \leftarrow ((\mathbf{b}_{(k)}\mathbf{y})^4 - 3) - \gamma_{(k-1)}
15:
                 end while
16:
                 x_i = \mathbf{b}^T \mathbf{y}
17:
18:
           end for
     end procedure
```

C.1.5 ICA for sparse signal recovery

ICA is widely used within sparse signal recovery. When ICA is applied to a measurement vector $\mathbf{y} \in \mathbb{R}^M$ it is possible to separate the mixed signal into M or less independent components. However, by assuming that the independent components make a k-sparse signal it is possible to apply ICA within sparse signal recovery of cases where M < N and $k \le M$.

To apply ICA to such cases the independent components are obtained by the pseudo-inverse solution

$$\hat{\mathbf{x}} = \mathbf{A}_S^\dagger \mathbf{y}$$

where \mathbf{A}_S is derived from the dictionary matrix \mathbf{A} by containing only the columns associated with the non-zero entries of \mathbf{x} , specified by the support set S, cf. appendix A.1.

C.2 Fixed-Point Algorithm - FastICA

An advantage of gradient algorithms is the possibility of fast adoption in non-stationary environments due the use of all input, \mathbf{y} , at once. A disadvantage of the gradient algorithm is the resulting slow convergence, depending on the choice of

 γ for which a bad choice in practise can disable convergence. A fixed-point iteration algorithm to maximise the non-Gaussianity is an alternative that could be used.

Consider the gradient step derived in section C.1.4. In the fixed point iteration the sequence of γ is omitted and replaced by a constant. This builds upon the fact that for a stable point of the gradient algorithm the gradient must point in the direction of **b**, hence be equal to **b**. In this case adding the gradient to **b** does not change the direction and convergence is achieved. Letting the gradient given in (C.4) be equal to **b** and considering the same simplifications again suggests the new update step as [19, p. 179]

$$\mathbf{b} \leftarrow \mathbb{E}[\mathbf{y}(\mathbf{b}^T\mathbf{y})^3] - 3\mathbf{b}.$$

After the fixed point iteration **b** is again divided by its norm to withhold the constraint $\|\mathbf{b}\| = 1$. Instead of γ the fixed-point algorithm compute **b** directly from previous **b**.

The fixed-point algorithm is referred to as FastICA. The algorithm has shown to converge fast and reliably, when the current and previous **b** point in the same direction [19, p. 179].

C.2.1 Negentropy

An alternative measure of non-Gaussianity is the negentropy, which is based on the differential entropy. The differential entropy H of a random vector \mathbf{y} with density $p_{y}(\boldsymbol{\eta})$ is defined as

$$H(\mathbf{y}) = -\int p_y(\boldsymbol{\eta}) \log(p_y(\boldsymbol{\eta})) d\boldsymbol{\eta}.$$

The entropy describes the information that a random variable gives. The more unpredictable and unstructured a random variable is higher is the entropy, e.g. Gaussian random variables have a high entropy, in fact they have the highest entropy among the random variables of the same variance [19, p. 182].

Negentropy is a normalised version of the differential entropy such that the measure of non-Gaussianity is zero when the random variable is Gaussian and non-negative otherwise. The negentropy J of a random vector \mathbf{y} is defined as

$$J(\mathbf{y}) = H(\mathbf{y}_{\text{gauss}}) - H(\mathbf{y}),$$

with $\mathbf{y}_{\text{gauss}}$ being a Gaussian random variable of the same covariance and correlation as \mathbf{y} [19, p. 182]. As the kurtosis is sensitive for outliers, the negentropy is instead difficult to compute computationally as the negentropy require a estimate of the pdf. As such an approximation of the negentropy is needed. To approximate the negentropy it is common to use the higher order comulants including the kurtosis.

The following approximation of the scalar case is stated without further elaboration, the derivation can be found in [19, p. 183].

$$J(\mathbf{y}) \approx \frac{1}{12} \mathbb{E}[y^3]^2 \frac{1}{48} \text{kurt}(y)^2.$$

C.2.2 Fixed-Point Algorithm with Negentropy

Maximization of negentropy by use of the fixed-point algorithm is now presented, for derivation of the fixed point iteration see [19, p. 188]. Algorithm 5 show Fast ICA using negentropy, this is the algorithm which is applied in the thesis for comparison with the source recovery methods which are tested in this thesis.

Algorithm 5 Fast ICA – with negentropy

```
1: procedure Pre-Processing(y)
            Center measurements \mathbf{y} \leftarrow \mathbf{y} - \bar{\mathbf{y}}
            Whitening \mathbf{y} \leftarrow \mathbf{y}_{white}
 3:
 4: end procedure
 5:
 6: procedure FASTICA(y)
            k = 0
 7:
            Initialise random vector \mathbf{b}_{(k)}
                                                                                                                              ▶ unit norm
 8:
            for j \leftarrow 1, 2, \dots, N do
 9:
                  while convergance critia not meet do
10:
                        k = k + 1
11:
                       \mathbf{b}_{(k)} \leftarrow \mathbb{E}[\mathbf{y}(\mathbf{b}_{(k-1)}^T \mathbf{y})] - \mathbb{E}[g'(\mathbf{b}_{(k-1)}^T \mathbf{y})] \mathbf{b}_{(k-1)} \qquad \triangleright g \text{ cf. [19, p. 190]}
\mathbf{b}_{(k)} \leftarrow \mathbf{b}_{(k-1)} / ||\mathbf{b}_{(k-1)}||
12:
13:
                  end while
14:
                  x_i = \mathbf{b}^T \mathbf{y}
15:
16:
            end for
17: end procedure
```

C.3 Verification of fast ICA on synthetic data

The purpose of this section is to verify the fast ICA algorithm which is used in this thesis. By this verification the purpose is to justify the ICA algorithm as a reference point with respect to performance of the developed main algorithm.

The fast ICA algorithm is tested on synthetic data simulated as described in section 6.2. Consider the following linear system, which makes a model of EEG measurements.

where $\mathbf{Y}^{M\times L}$, $\mathbf{A}^{M\times N}$ and $\mathbf{X}^{N\times L}$. It is expected that the fast ICA algorithm manage to solve the linear system for \mathbf{X} and \mathbf{A} given only the measurements \mathbf{Y} , in the case where M=N.

The fast ICA algorithm is applied to \mathbf{Y} and returns the estimates $\hat{\mathbf{X}}_{ICA}$ and $\hat{\mathbf{A}}_{ICA}$. When using the fast ICA algorithm the output $\hat{\mathbf{X}}_{ICA}$ do not correspond one to one with the true source signal, which become an issue when the estimation error is measured by the mean squared error (MSE) cf. subsection 6.2.3. The fast ICA algorithm is invariant towards the amplitude and phase of the signal and furthermore the rows are not necessarily place the original location. In order to get a valid MSE measure of the estimate, a function is defined to fit the estimate to the true source signal \mathbf{X} . The function manage to pair the rows and change the phase, such that the total MSE is minimized. Furthermore each row of the estimate is scaled by the relationship between the maximum value of the true row and the estimated row. From empirical observations only the phase shift performed by multiplying with (-1) has shown necessary, hence it is easily applied to the fitting function. When the fitting function is applied two the estimate, the full potential of the fast ICA algorithm is considered reached.

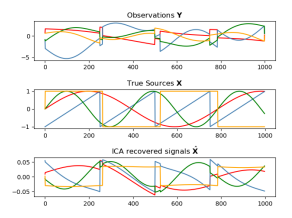
Figure C.1 illustrates $\hat{\mathbf{X}}_{ICA}$, without use of the fitting function, resulting from the fast ICA algorithm applied to a simulated deterministic data set \mathbf{Y} specified by M = N = k = 4 and L = 1000. In the figure \mathbf{Y} , \mathbf{X} and $\hat{\mathbf{X}}_{ICA}$ are plotted separately and it is clear to see the invariance towards amplitude and phase. The MSE from the original $\hat{\mathbf{X}}_{ICA}$ becomes

$$MSE(\mathbf{X}, \hat{\mathbf{X}}_{ICA}) = 0.608.$$

In figure C.2 the fitting function has been applied to $\hat{\mathbf{X}}_{ICA}$. Each row of the fitted estimate is now plotted with the corresponding row of the true source signals. The resulting MSE becomes

$$MSE(\mathbf{X}, \hat{\mathbf{X}}_{ICA}) = 0.046.$$

This is an essential change from the first measured MSE, and it is considered to provide a more valid measure of the estimate. From the visualisation and the corresponding MSE it is found that the fast ICA algorithm manage to estimate the source signals of the deterministic data set with a sufficiently small error.



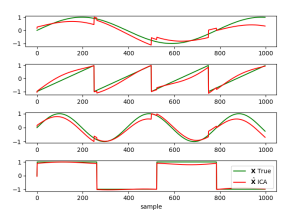


Figure C.1: Plot of simulated deterministic observations Y, specified by M = N = k = 4 and L = 1000. Corresponding plot of the true \mathbf{X} and the estimated $\hat{\mathbf{X}}$ by ICA.

Figure C.2: Direct comparison of the true \mathbf{X} and $\hat{\mathbf{X}}_{ICA}$ after applying the fitting function

A similar test is now performed on a stochastic data set \mathbf{Y} , cf. section 6.2.2, again specified by M=N=k=4 and L=1000. Figure C.3 show the comparison of the fitted $\hat{\mathbf{X}}_{ICA}$ and the true source signals \mathbf{X} . Note that only the first 100 samples are plotted for easier visualization. The resulting MSE becomes:

$$MSE(\mathbf{X}, \hat{\mathbf{X}}_{ICA}) = 0.037.$$

Again the the MSE is considered sufficiently small and by that the fast ICA is considered verified with respect to solving a linear system with M = N.

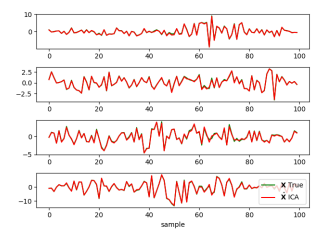


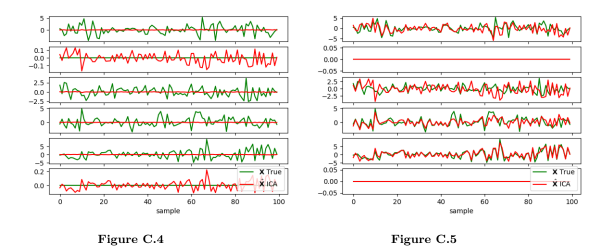
Figure C.3: ICA applied to stochastic simulated observations Y. Direct comparison of the true X and $\hat{\mathbf{X}}_{ICA}$ after applying the fitting function.

Consider now the case where $k \leq N = M$, that is the sources signal matrix has k non-zeros rows. The fast ICA algorithm is now applied to a stochastic data set **Y**

specified by N = M = 6, k = 4 and L = 1000. Figure C.4 and C.5 show the comparison of the resulting $\hat{\mathbf{X}}_{ICA}$ and the true \mathbf{X} before and after the application of the fitting function, respectively. The resulting MSE becomes:

$$MSE(\mathbf{X}, \hat{\mathbf{X}}_{ICA}) = 1.784.$$

It is seen from figure C.5 that the fast ICA algorithm manage to detect the zero rows of \mathbf{X} . Without further test, this indicates the possibility of estimating k from the fast ICA algorithm.



With these tests the quality of the fast ICA algorithm has been verified. As such the fast ICA algorithm can be used as a reference, when applied to real EEG data. It is further established that $k \leq M$ can be estimated by fast ICA.

Remember though that the ICA estimate is conditioned under $k \leq N = M$, however this condition is not necessarily withhold for real EEG measurements as the true N is always unknown.

Appendix D

Python Scripts

The following list contain descriptions of the essential Python scripts which have been used through this thesis.

- 1. Data_Simulation.py: Contain functions for simulation of deterministic and random synthetic data, based on manual specification of each system.
- 2. Data_EEG.py: Contain functions for import of EEG measurments, possible reduction and segmentation.
- 3. ICA_Fast.py: Contain function to perform ICA.
- 4. Main_Algorithm.py: In this script the main algorithm composed by a compilation of the necessary modules. Remark, by letting fix = True Cov-DL is overwritten by $\hat{\mathbf{A}}_{fix}$.
 - Cov_DL.py: Contain the function to perform Cov-DL
 - M_SBL.py: Contain the function to performM-SBL
- 5. Test_Synthetic_data.py: Compilation of necessary modules to apply the main algorithm to on synthetic data and generate output, cf. chapter 6.
- 6. Test_EEG.py: Compilation of necessary moduels to apply ICA and the main algorithm on EEG measurements, for the 3 different cases, cf. section ??.
- 7. Test_AlphaFrequency.py: Contain functions and corresponding compilation to perform alpha wave analysis, cf. section 7.4.
- 8. Test_k_Estimation.py: Contain functions and corresponding compilation generate estimation of k, cf. chapter 8

The Python scripts are all available directly at https://inset_link.git. Besides the script, the folder contains the EEG measurements and a folder for which generated plots are allocated, is present.