Analysis and visualization of phase synchronization in EEG signals during cognitive tasks

TESIS

Que para obtener el Grado de

Doctorado en Ciencias de la Computación

PRESENTA:

Francisco Alfonso Alba Cadena

Directores de Tesis:

Dr. José Luis Marroquín Zaleta
Dra. Thalía Harmony Baillet

GUANAJUATO, GTO.

Analysis and visualization of phase synchronization in EEG signals during cognitive tasks

TESIS

Que para obtener el Grado de

Doctorado en Ciencias de la Computación

PRESENTA:

Francisco Alfonso Alba Cadena

Comité de Evaluación:

Dr. Johan Jozef Lode Van Horebeek
(Presidente)

Dr. Mariano José Juan Rivera Meraz
(Secretario)

Dr. Rogelio Hasimoto Beltrán
(Vocal)

Dr. Rogelio Ramos Quiroga
(Vocal)

Dra. Thalía Harmony Baillet
(Vocal y co-directora de Tesis)

Director de Tesis:

Dr. José Luis Marroquín Zaleta

GUANAJUATO, GTO.  \hspace{2cm} MARZO DE 2007.
Abstract

In this thesis we propose a new methodology for the study of phase synchronization in electroencephalographic signals. This methodology attempts to overcome some of the drawbacks found in the procedures typically used in the literature by using (1) sinusoidal quadrature filters, which give a more reliable response when tuned at low frequencies, (2) instantaneous in-phase synchrony measures, (3) a Bayesian classification approach to determine the significance of the synchrony changes with respect to a neutral condition, and (4) a time-frequency-topography (TFT) visualization system which can display the results with high spatial, temporal, and frequency resolutions. All these techniques are implemented in a software program which has already been used by neuroscientists to analyze various psychophysiological experiments. We have also tested our methodology with various of the synchrony measures proposed in the literature, and performed a comparative study which highlights the advantages and disadvantages of each one. Finally, we propose a simple mathematical model for the narrow-band EEG signal observed at each electrode, which allows one to explain and simulate different synchronization patterns that can be observed with real EEG data.
Contents

1 Introduction 4
  1.1 Introduction to EEG ................................. 4
  1.2 EEG applications ................................. 5
  1.3 Analysis of EEG signals ........................... 6
    1.3.1 Evoked potentials ............................ 6
    1.3.2 Power analysis ............................... 6
    1.3.3 EEG inverse problem .......................... 7
    1.3.4 Synchrony analysis ............................ 7
  1.4 EEG rhythms ....................................... 7
  1.5 Reference electrode ............................... 8
  1.6 Volume conduction .................................. 8
  1.7 Goals ............................................. 11
  1.8 Thesis organization ............................... 12
  1.9 Experimental data ................................. 12
    1.9.1 Figures and Words experiments ................. 13
    1.9.2 3-digit and 5-digit Sternberg experiments .... 13
    1.9.3 Letters experiment ............................ 14

2 Methods and measures for the study of EEG synchrony 15
  2.1 Introduction ...................................... 15
  2.2 Time-frequency decomposition ....................... 16
    2.2.1 Local windowed Fourier transform ............... 16
    2.2.2 Gabor filters .................................. 17
    2.2.3 Sinusoidal quadrature filters .................... 19
    2.2.4 Wavelet analysis ................................ 19
  2.3 Long-range synchrony measures ...................... 20
    2.3.1 Coherence ..................................... 21
    2.3.2 Phase Locking Statistic ....................... 23
    2.3.3 Single-Trial Phase Locking Statistic .......... 25
    2.3.4 Generalized Synchronization .................... 26
    2.3.5 Mutual Information ............................ 27
  2.4 Statistical significance analysis ................... 28
    2.4.1 Choosing the null population .................... 28
    2.4.2 Significance estimation: normal distribution .... 29
<table>
<thead>
<tr>
<th>Section</th>
<th>Title</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.4.2</td>
<td>Anti-phase synchrony</td>
<td>87</td>
</tr>
<tr>
<td>6.4.3</td>
<td>Synchrony at multiple frequencies</td>
<td>90</td>
</tr>
</tbody>
</table>
Chapter 1

Introduction

1.1 Introduction to EEG

Electroencephalography (EEG) is a neuroimaging technique that consists of voltage measurements recorded by one or more electrodes placed on the scalp (non-invasive EEG), or within the cortex (intracranial EEG). The first person to obtain an EEG recording from a human brain was Hans Berger, in the 1920’s. He was also the first to observe the cyclic nature of the EEG and the relation between different states of consciousness (e.g. sleep, attention processes, wakefulness, etc.) and specific EEG wave patterns, especially in the frequency range from 8 to 12 Hz (alpha range) [Berger, 1929]. Figure 1.1 shows one of the EEG recordings obtained by Berger.

Although there are some recent intracranial studies (e.g. [Lachaux et al., 2000a]), most human EEG recordings are non-invasive: the electrodes are mounted in a cap, which is put in the subject’s head. The electrodes are usually systematically distributed across the scalp surface; the most widely used distribution corresponds to the 10-20 international electrode placement system (shown in Figure 1.2). This system standardizes the electrode positions and nomenclature, and also allows researchers to compare their analysis and visualization.

Figure 1.1: One of the first human electroencephalograms (Berger, circa 1929): the top trace is the EEG recording and the bottom trace is a 10 Hz reference signal.
1.2 EEG applications

There are numerous applications of EEG in modern medicine. Some examples are:

- **Study of the properties of neural networks** - By properly designing an experiment and adequate analysis of the EEG recordings, scientists may obtain insights about neural activity and/or prove certain hypotheses which are specifically related to the task performed by the subjects.

- **Diagnostic** - EEG is used in brain-mapping for localizing the areas of epileptic activity prior to operation.

- **Therapy** - Neurofeedback consists of the self-modification of electrophysiological brain patterns. It may be used to treat psychological and neurological disorders, and also to learn mental relaxation.

We will focus on the first application, in which a psychophysiological experiment is designed, performed, and analyzed, in order to study the brain activity during the experiment. Most EEG experiments involve multiple subjects, and each subject performs the experiment multiple times. Each repetition of the experiment is called a trial, and the recorded data can be seen as a set of voltage time series $V_{j,e}(t)$ where $j$ is the trial number, $e$ is the index of the recording channel or electrode (representing the spatial position of the sensors), and $t$ is the time index.
A wide range of psychophysiological experiments follow this scheme: each subject is presented with a stimulus (e.g., a certain image in a screen, a sound or a word, etc.) and is instructed to respond in some way (e.g., press a button) or perform a certain task (e.g., add two numbers, memorize a set, etc.). EEG is recorded during the experiment, one segment per trial. Each segment starts before the stimulus presentation and finishes after the subject has responded. Thus each segment is commonly split in two parts: the pre-stimulus segment corresponding to the recording before the stimulus onset, and the post-stimulus segment, which contains all activity after the stimulus. In these experiments, one usually performs a statistical comparison between both pre and post-stimulus segments, in order to detect significant changes in the neural activity which may be related to the task performed by the subjects.

1.3 Analysis of EEG signals

In order to obtain information about the neural events taking place during the experiment, different types of analysis may be performed with the EEG data. These usually involve time-frequency decomposition techniques and statistical tools. Here we describe the most common types of analysis found in the literature.

1.3.1 Evoked potentials

Evoked activity corresponds to EEG changes which are time-locked and phase-locked to the stimulus [Pfurtscheller and Lopes da Silva, 1999]. This activity is usually obtained, for each recording channel, by averaging the recorded potentials across all trials:

\[
EP_e(t) = \frac{1}{N_t} \sum_{j=1}^{N_t} V_{j,e}(t). \tag{1.1}
\]

Here \(N_t\) is the number of trials, and \(EP_e\) represents the evoked activity at electrode \(e\). These averaged signals may show significative peaks (called event-related potentials or ERP’s) which may be related to the task. ERP’s are named according to their latency with respect to the stimulus onset (in milliseconds), and the letter N or P indicating its sign. For example, P300 denotes a positive evoked potential taking place around 300 ms after the stimulus.

1.3.2 Power analysis

One of the most common tools used for EEG analysis consists on a spectral power decomposition at the frequency bands of interest. It is known that certain frequency bands are related to different states of attention and consciousness (see [Niedermeyer, 2004], Chapter 3); for example, the power ratio between certain pairs of bands may be used for neurofeedback therapy. Time-frequency
power analysis is also useful to detect event-related induced activity, which is time-locked but not phase-locked to the stimulus. The increases or decreases of power in specific frequency bands may be considered to be due to an increase or decrease, respectively, in the synchrony of the underlying neuronal populations [Pfurtscheller and Lopes da Silva, 1999], and thus are accordingly called “event-related synchronization” (ERS) and “event-related desynchronization” (ERD). Other works have also shown that changes in power may be related to cognitive and attentional processes (see [Harmony et al., 2004; Marroquin et al., 2004]).

1.3.3 EEG inverse problem

The inverse problem in EEG consists on localizing the current sources which are located in the brain, and which give origin to the potentials recorded by the sensors. This is an ill-posed problem and the solution is not unique; thus one must impose restraints to find a particular solution that suits a specific problem (a good review on some inverse problem methods can be found in [Pascual-Marqui, 1999]).

1.3.4 Synchrony analysis

EEG synchrony studies the dynamical connectivity patterns across neural networks, and their function as integration mechanisms. Synchrony can be studied at two levels: first, from a local point of view, where synchronization and desynchronization events between networks within a small area (separated by less than ∼1 cm) may be reflected as power changes or bursts in the corresponding EEG channel [Varela et al., 2001]; and second, there is evidence for long-range synchronization between distant networks (not necessarily contiguous), which is reflected as some form of correlation or phase-locking between the signals from different sensors [Bressler et al., 1993; Rodriguez et al., 1999; Hoechstetter et al., 2004; Mizuhara et al., 2005].

1.4 EEG rhythms

Since the introduction of the EEG by Berger in the 1920’s, researchers have recognized, categorized, and described the cyclic waveforms of EEG recordings (called EEG rhythms). These rhythms have been classified according to their frequency as: delta (1-4 Hz), theta (4-7 Hz), alpha (7-14 Hz), beta (14-30 Hz) and gamma (30 Hz and upwards). A thorough discussion of the neurophysiological origin of each of these rhythms can be found in [Niedermeyer, 2004] (Chapter 3).

Although it is very common to focus a particular analysis in one or more of these rhythms, it is recommended to perform the time-frequency decomposition in narrow frequency bands in order to obtain meaningful values of instantaneous amplitude and phase.
1.5 Reference electrode

The EEG measurements in a specific channel actually represent the difference of electric potentials at two electrode sites: the electrode corresponding to that channel (target electrode), and a reference electrode [Hagemann et al., 2001]. Electrical activity at either of these electrodes will be reflected in the EEG. Because of this, different placements of the reference electrode may result in very different derivations for each target electrode. Ideally, one would place the reference on a site with very low or null electrical activity. In practice, most recordings use a reference site placed on or near the head (cephalic reference), such as the neck, chin, or linked earlobes. It is also possible to use a non-cephalic reference; however, this may result in recordings largely contaminated by electrocardiogram (EKG) artifacts [Stephenson and Gibbs, 1951].

The reference problem is important in synchrony analysis because it may introduce spurious correlations between different EEG channels. However, in a cognitive task experiment, one may assume that any activity at the reference electrode is not related to the task, and thus the reference signals for different trials may be assumed to be uncorrelated, and its effect may be reduced after averaging across trials.

There are a few methods to transform the raw EEG signals into a new data set which further reduces the reference effects. Two of them are (1) current source density (CSD) estimation, which will be discussed in the following section, and (2) transformation to the average reference (AR). The AR data set is obtained by estimating the “average electrode” signal (which is the average signal of all lead signals) and subtracting it from each electrode signal to obtain a new derivation. This has the effect of re-referencing all the signals to a new virtual reference site whose activity is precisely the average of the theoretical reference-free lead signals. The problem is that, in practice, this activity is not only far from null but it is also biased towards the center the covered region, and it may also reflect changes related to the stimulus presentation in a cognitive experiment (e.g. evoked potentials may induce high activity at the virtual reference site), thus the effect of this virtual reference may not be canceled out by averaging across trials [Junghöfer et al., 1999].

An interesting discussion of different derivation methods (including linked ears, average reference, and CSD estimation) can be found in [Hagemann et al., 2001]. Hagemann’s findings suggest that each method has its own limitations and some of them may be more appropriate for a specific type of analysis than the others.

1.6 Volume conduction

Another known problem is caused by the conductive properties of the cortex, skull, and scalp, which smear the potentials across the surface and reduces the
spatial resolution of the EEG [Nunez, 1995]. In 1975, Hjorth proposed a mathematical procedure to estimate the “source activity at the surface level of the scalp” [Hjorth, 1975], which is known as scalp current density (SCD) or current source density (CSD). This procedure effectively reduces volume conductor effects and yields a new set of signals which are more localized (i.e. have better spatial resolution) than the raw potentials. Moreover, CSD is not sensitive to the reference potential.

The idea behind CSD relies on estimating the current density field \( \bar{J} \) from Ohm’s law:

\[
\bar{J} = \sigma E,
\]

where \( \sigma \) is the conductivity and \( E \) is the electric field. Given a surface distribution of potentials \( \Phi \) we may calculate the electric field as \( E = -\nabla \Phi \), thus

\[
\bar{J} = -\sigma \nabla \Phi.
\]

Finally, the divergence of a vector field represents the source intensity of the field. We apply this to the current density field \( \bar{J} \) to obtain the current source density \( J \):

\[
J = \nabla \cdot \bar{J} = -\sigma \nabla^2 \Phi.
\]

According to Equation 1.4, CSD is negatively proportional to the surface Laplacian (SL) of the potentials. In practice, one is usually interested in relative changes of some measure with respect to the pre-stimulus, and the factor \( -\sigma \) is canceled; thus one may work only with the SL.

During the following decade, Hjorth’s procedure was refined to obtain faster and more accurate results (see [Pascual-Marqui et al., 1988], [Perrin et al., 1988, 1989]; [Law et al., 1993]). Basically, the SL is estimated by interpolating the potentials analytically for each time \( t \), and then taking the second spatial derivatives of the interpolating function. Presently, the most used method is the one introduced by Perrin et al. in 1988, which approximates the SL by using spherical splines. More recently, Mizuhara et al. used CSD estimates to study long-range phase synchronization in a cognitive task [Mizuhara et al., 2005] and combined it with fMRI to determine which networks were being integrated during phase synchronization stages. (Functional MRI or fMRI is 3D imaging technique which is used to locate which parts of the brain are activated by showing regions with increased blood flow).

Nunez et al. [1995; 1997; 2000] have developed an extensive work on the effect of volume conduction in EEG coherence. Using simulated data from a 3 concentric spheres model, they estimated correlation coefficients for pairs of uncorrelated cortical sources using different EEG references (Cz, neck, linked ears, and average reference), cortical imaging, and Spline Laplacians ([Perrin et al., 1989, 1990; Law and Nunez, 1993]). In all cases, except Spline Laplacians, spurious high correlations were observed for short inter-electrode distances (4-8 cm) [Nunez et al., 1997]. However, since the Surface Laplacian acts as a
bandpass filter on the raw potentials, true coherence with low spatial frequency (i.e., relatively smooth across the surface) may be underestimated. Because of this, Nunez suggests to use the SL to complement, rather than replace, the raw EEG potentials: correlations may be observed at different spatial scales [Nunez et al., 1997, 2000] (just as correlations between customs of people from two different cities may differ from correlations between two families, one from each city). Therefore, raw potential, Surface Laplacian, and cortical image coherency may, in general, represent correlations at different spatial scales of cortical dynamics.

Unfortunately, there is a problem when measuring phase synchronization from SL signals: since the potentials are a smooth function of the position in the surface (partly because volume conductor effects), the Surface Laplacian, which is the second spatial derivative of the potentials, will have a very small magnitude. This means that the phase of the SL signals will be very sensitive to noise, and also to the interpolation method used to estimate the SL. Hagemann et al. [2001] also note this problem in his comparison of different reference derivations. According to [Junghöfer et al., 1999], current density measures with 32 or less channels will be unstable and inaccurate, and even for high spatial sampling density, the second derivative is much more vulnerable to noise than the scalp potentials.

An alternate solution to the volume conductor problem consists on estimating the current sources that give origin to the surface potentials. This usually involves placing various dipoles in the areas of interest and solving the inverse problem (with some necessary constraints) to find the source signals. This approach has been used by David et al. [2001], and Hoechstetter et al. [2004]. In David’s method, the inverse problem is restrained by penalizing solutions which have sources with large amplitude that do not contribute to the frequency band of interest; the amount of penalization is locally determined (i.e. differs from source to source) and depends on the amount of power outside the frequency band of interest: if the signal in a given source has most of its power outside the frequency band of interest, that source will be highly penalized. Synchronization measures are usually calculated for each pair of sources, thus it is desirable to have as few sources as possible while maintaining an adequate spatial resolution. In order to minimize the number of reconstructed dipoles, David et al. use the following iterative approach: first, start with a large number of dipoles distributed uniformly across the whole cortex (with radial orientations) and solve the inverse problem for these sources. Then, the best dipoles are selected (according to a fitness measure, which is also based in the signal power outside the frequency band of interest) and the process is repeated for the selected dipoles until their fitness is higher than some threshold. Once the sources have been determined, the Single-Trial Phase-Locking Statistic (STPLS - [Lachaux et al., 2000b]), which we will describe in the next chapter, is estimated for each source pair.

One problem with the method described above is that it heavily relies on the spectral properties of the signals estimated for the full post-stimulus segment,
but does not account for possible changes of power during the course of an experiment (EEG signals are hardly static). Moreover, to analyze different frequency bands, one must solve the inverse problem and obtain a different set of sources for each band, which may provide a difficult interpretation of the results across bands.

The approach used by Hoechstetter is much simpler: the sources are placed in sites of interest which can be either pre-specified, or obtained from evoked data. Also, additional dipoles may be placed to model background activity (e.g. eye movement activity). These dipoles function as virtual sources within the brain. Later, a coherence measure (see [Gardner, 1992]) is estimated for each pair of source signals.

The problem, however, is that this method assumes that the modeled sources account for all activity recorded in the sensors and, unlike the adaptive selection approach used by David et al., this method does not attempt to find the sources whose activity may be related to the synchronous processes. Therefore, if the sources are not adequately distributed, the signals obtained for some of the dipoles may in fact represent the activity in large, overlapped cortical regions, which would defeat the purpose of source localization. Hoechstetter suggests fitting the sources to evoked activity; however, the relation between long-range synchrony and evoked activity is not yet clear.

1.7 Goals

The main goal of this work is to present a robust methodology to detect, analyze, and visualize significant changes in patterns of dynamic connectivity in both a wide spatial level (long range synchrony), and a local level (power analysis and evoked potentials). These changes may happen within a narrow frequency band and last only few dozens of milliseconds; thus it is important to preserve the time and frequency resolutions. It is also desirable that the techniques described here may be used with different synchrony measures (e.g. coherence, mutual information, etc - see Chapter 2 for more details).

While we focus in electroencephalographic signals in this work, it is intended that the most important contributions of this methodology hold for several types of neurophysiological signals, including: intracranial EEG, current sources, current density, and MEG.

The final procedure will be implemented as a Windows program with a straightforward interface and various navigating tools in order to easily identify and focus on the regions of interest. The program will

1. Allow the user to input raw EEG data in a standard file format
2. Preprocess and analyze the data to estimate and classify synchrony changes
3. Display the results in various types of graphs
4. Allow the user to save the results in a file (for further analysis with other programs)

As an example of application, the program described above will be used by neuroscientists to study the connectivity patterns in real EEG experiments from a neurophysiological point of view. We expect at least one paper to be published from these studies. Another paper describing the methodology (from a technical point of view) is already in print.

1.8 Thesis organization

The contents of this thesis are organized as follows:

- **Chapter 2** presents a summary of the most common methods and techniques used for EEG synchrony analysis. It also describes the most popular long-range synchrony measures found in the literature.

- **Chapter 3** presents a detailed description of the proposed methodology and synchrony measures.

- In **Chapter 4** we discuss different aspects of the synchrony measures presented in Chapter 2 and the ones proposed in Chapter 3, from a comparative point of view.

- **Chapter 5** focuses on a mathematical model with which one can simulate (and explain to some extent) different phenomena observed during the analysis of various real experiments.

- **Chapter 6** summarizes the main contributions of this work, and also presents the results of the neurophysiological analysis of real experiments. Ideas for future work are also presented here.

- **Appendix ??** presents the resulting graphs and results for additional experiments.

- **Appendix ??** describes the publications and presentations originated from this thesis.

- **Appendix ??** contains the user manual for our EEG analysis software.

1.9 Experimental data

During the course of our work, we have tested the procedures using data from various EEG experiments. Here we describe each of them.
1.9.1 Figures and Words experiments

This experiments are designed for the study of categorization tasks [Harmony et al., 2001]. For the Figures experiment, white-line figures on a black background were presented to each subject. The subjects were instructed to press a button if the figure corresponded to an animal whose name started with a consonant, and another button if the figure did not correspond to an animal and the name of the figure started with a consonant. If the name started with a vowel, the subject was instructed not to respond.

The Words experiment follows a similar paradigm, with the exception that words instead of figures are presented on the screen. The subjects were instructed to press one button if the word corresponded to an animal and starts with a consonant, and another button if the word did not correspond to an animal and started with a consonant. Otherwise, the subject was instructed not to respond.

In both experiments, the subjects were 18 normal children (8 to 10 years old, 9 females), all right handed with normal neurological examination. EEG was recorded with reference to linked ears from Fp1, Fp2, F3, F4, C3, C4, P3, P4, O1, O2, F7, F8, T3, T4, T5, T6, Fz, Cz, Pz, and Oz of the 10/20 system. EOG was recorded from a supraorbital electrode and from an electrode on the external canthus of the right eye. The amplifier bandwidth was set between 0.5 and 30 Hz. EEG was sampled every 5ms using a MEDICID 3E system and stored on a hard disk for further analysis. Sampling was done every 5 ms during a time segment from 1280 ms before the stimulus to 1500 ms after its onset. Each trial was visually edited and only those corresponding to correct responses and with no artifacts were analyzed. Subjects were seated in a comfortable chair in front of the videomonitor. Stimuli were delivered by a MINDTRACER system synchronized to the MEDICID 3E acquisition system.

1.9.2 3-digit and 5-digit Sternberg experiments

In this experiment [Harmony et al., 2004], a visual warning stimulus (lasting 300 ms) is presented at the start of each trial. After two seconds, a set of three or five digits (memory set) is presented for 1500 ms. Two seconds later, a single digit (probe stimulus) is displayed for 300 ms. The subject must respond with one button if the probe was in the memory set, and with another button if it was not. In 50% of the trials, the number belonged to the memory set. The EEG was sampled each 5 ms with reference to linked ears from Fp1, Fp2, F3, F4, C3, C4, P3, P4, O1, O2, F7, F8, T3, T4, T5, T6, Fz, Cz, Pz, and Oz. Segments of 1280 ms were selected immediately before the memory set was presented (pre-segment) and immediately before the presentation of the probe stimulus (post-segment). Only correct responses and artifact-free segments were analyzed.
1.9.3 Letters experiment

This experiment corresponds to a Go/NoGo task designed to study the inhibition of the motor response. A series of uppercase letters are shown, one at a time, on the screen. The interval between the presentation of one letter and the next one is two seconds. The subject is instructed to respond with a button only if a 'X' that has been preceded by an 'O' appears. This is the Go condition. Any letter different than 'X' which has been preceded by an 'O' accounts for the NoGo condition, as it may originate the inhibition of the motor response. EEG was recorded each 5 ms from Fp1, Fp2, F3, F4, C3, C4, P3, P4, O1, O2, F7, F8, T3, T4, T5, T6, Fz, Cz, and Pz. Each segment was selected from 1 s immediately before the presentation of each letter (pre-segment) to 1560 ms immediately after (post-segment). The analysis of this experiment using the techniques described below has been submitted for publication [Harmony et al., 2006].
Chapter 2

Methods and measures for the study of EEG synchrony

2.1 Introduction

It is thought that during the execution of a relatively complex task, specialized (and possibly distant) areas of the brain interact together by means of strong reciprocal connections, forming what is called a neural assembly [David et al., 2003a]. The mechanisms involved in this large-scale integration are still largely unknown; however, one of the most plausible candidates is the formation of dynamical links that are reflected as some form of synchronization over different frequency bands [Varela et al., 2001] (in particular, Varela favors phase-synchronization as the most plausible mechanism).

According to Varela et al., synchronization of neural assemblies is a process that spans multiple spatial and temporal scales. In this sense, the most adequate level of synchrony analysis would be at single neuron level: in the timescale of cognitive events (typically hundreds of ms), a single neuron may fire only a few spikes, which may not be enough to activate a target neuron. However, if these spikes coincide in time with those fired by additional neurons, the target neuron may be activated. Unfortunately, it is impossible to study neural interactions at this level with the present EEG scalp sensors. On a coarser level, it is possible to record the activity of a local neural group (within a few millimeters) using intracranial EEG sensors: the recorded Local Field Potentials (LFP’s) blur the contributions of the participating neurons, but at the same time highlight their common actions. In this case, a time-frequency power analysis may yield information about integration within the local network.

Finally, there are also studies of long-range synchronization, where the integrated networks are separated by ~1 cm or more. This involves reciprocal, poly-synaptic communication paths across the networks, which result in transmission delays of at least 8 ms. It is known that the brain areas are massively
and reciprocally connected, and that such a connectivity engenders the emergence of correlations in the brain activity of the different areas [David et al., 2003a]. Numerous methods have been used to study these correlations. Most of these follow a particular scheme: (1) (optionally) pre-process the EEG signals to reduce reference and volume conduction effects, (2) perform a time-frequency decomposition of the signals, (3) estimate a synchronization measure between pairs of lead signals of interest, and (4) perform a statistical test to determine the significance of the measured synchrony with respect to the baseline. Certain aspects of (1) have already been discussed in Sections 1.5 and 1.6. We will also propose a projection method to reduce volume conduction in Section 3.7. The rest of this chapter will deal with steps (2) through (4).

2.2 Time-frequency decomposition

Electroencephalographic recordings have a cyclic nature; therefore, it is very common to analyze them in the frequency domain rather than in the time domain. On the other hand, EEG is also very dynamic, and, in many cases such as in event-related analysis, the interest is precisely in detecting changes in the spectral properties of the signals, and in determining if those changes may be related to the task which is being studied. In these cases, it becomes necessary to perform a time-frequency (TF) decomposition of the signals from which one can extract instantaneous amplitude and phase information for each frequency band (the latter being particularly important for long-range synchrony analysis). In this section we present the most common tools used to achieve this.

2.2.1 Local windowed Fourier transform

The discrete version of the Fourier theorem states that a periodic discrete function \( f(t) \) can be expressed as a sum of pure complex tones as follows:

\[
f(t) = \frac{1}{N} \sum_{k=0}^{N-1} F(k) \exp \left[ \frac{2\pi i}{N} kt \right], \quad n = 0, \ldots, N-1, \tag{2.1}
\]

where \( N \) is the period of \( f(t) \) and \( F(k) \) are the Fourier coefficients given by

\[
F(k) = \sum_{t=0}^{N-1} f(t) \exp \left[ -\frac{2\pi i}{N} kt \right]. \tag{2.2}
\]

\( F \) is called the Fourier transform (FT) of \( f \), and it provides us with a description of the frequency content of the original signal. For example, the magnitude of \( F(k) \) indicates how much of the frequency \( 2\pi k/N \) is present in \( f \). In terms of EEG signals, this represents how strong is each rhythm in a given recording. One problem with the FT, however, is that it assumes the signals to be stationary (i.e. their frequency content does not change over time),
which may not be true for some signals such as EEG. The solution consists on estimating the Fourier transform “locally” within a narrow time window around $t_0$, and then just slide $t_0$ to obtain the Fourier coefficients $F(k,t_0)$ at each time. This is represented as

$$F(k,t_0) = \sum_{t=0}^{N-1} f(t)w(t-t_0) \exp \left[ -\frac{2\pi i}{N} kt \right], \quad (2.3)$$

where $w(t)$ is a real finite window function symmetric around zero. Generally, $w(t)$ decays smoothly as $|t|$ increases, so that only values relatively near $t_0$ have a significant weight when estimating $F(k,t_0)$. The width $\Delta t$ of the window has a direct effect in the time and frequency resolution of the local Fourier transform. Narrow windows will result in a higher time resolution, since the Fourier coefficients for $t_0$ will be less contaminated by values distant from $t_0$. However, the Heisenberg Uncertainty principle states that it is not possible to determine, with arbitrary precision, both the time and frequency localization of the components of a signal. Mathematically, the product of $\Delta t$ and the frequency width $\Delta \omega$ of the window function (that is, the width of the Fourier transform $W$ of $w$) has a lower bound. More specifically,

$$\Delta t \Delta \omega \geq \frac{1}{4\pi}, \quad (2.4)$$

with $\Delta t$ in seconds and $\Delta \omega$ in Hz. The lower bound in Equation 2.4 is reached when the window function is Gaussian, thereby providing the best compromise between time and frequency localization.

The local windowed Fourier transform is widely used in EEG research (for example, [Gross et al., 2001] and [Harmony et al., 2004]), however, it has the disadvantage of focusing only on a discrete set of frequencies given by $2\pi k S/M$, $k = 0, \ldots, M-1$, where $M$ is the length of the Fourier transform, and $S$ is the sampling rate (in Hz) of the analyzed signal. Even though the basic EEG rhythms lie on relatively wide bands, it is sometimes desirable to perform a time-frequency decomposition in narrow bands centered at arbitrary frequencies. This is especially important in phase-synchrony analysis, since the instantaneous phase of a signal, which can be obtained from the argument of the Fourier coefficients, only has a physical meaning for narrow-band signals. To overcome this, one can use a relatively large $M$ (length) for the Fourier transform, or resort to quadrature or wavelet filters.

### 2.2.2 Gabor filters

Gabor filters are quadrature band-pass filters which consist of a Gaussian kernel shifted in frequency to the tuning frequency $\omega$. The Gabor kernels are given by:

$$G_{\omega,\sigma}(t) = \frac{1}{Z} \exp \left[ -\frac{t^2}{2\sigma^2} \right] \exp [i\omega t], \quad (2.5)$$

17
Figure 2.1: True phase and estimated phase (using Gabor and sinusoidal quadrature filters) for a 1 Hz sinusoidal signal.

where $\sigma_t$ is the filter width (in time units), and $Z$ is a normalizing constant. The frequency response of a Gabor filter is a Gaussian centered at $\omega$ with a frequency width $\sigma_\omega$ given by:

$$\sigma_\omega = \frac{1}{4\pi \sigma_t};$$

in other words, Gaussian filters also reach the lower bound in the Heisenberg Uncertainty principle (Equation 2.4); because of this, they are also a common choice for EEG time-frequency decomposition (see [Marroquin et al., 2004; Schack et al., 2005]).

The complex filtered signal $F_\omega(t)$ is simply obtained as the convolution of the original signal $f$ and the filter kernel. The instantaneous amplitude and phase of the filter output can then be calculated from the magnitude and argument of $F_\omega(t)$.

A quadrature filter has the property of its frequency response being zero for all negative frequencies. For a Gabor filter, however, when the tuning frequency is relatively low, there is a chance of the filter having a strong response to negative frequencies. In this case, the quadrature property of the filter is destroyed, and the instantaneous magnitude and argument are no longer well-defined, which may lead to distorted phase estimates (see Figure 2.1).

This distortion can be easily characterized for sinusoidal signals. Given an input signal $x(t) = 2 \cos 2\pi ft$, the output $y(t)$ of a unit-gain bandpass filter tuned at $f$ Hz (obtained by convolving $x$ with the filter kernel) will have the form

$$y(t) = e^{i2\pi ft} + \alpha e^{-i2\pi ft},$$

where $\alpha \in [0,1)$ is the attenuation factor of the component with negative frequency. The argument $\Phi_y(t)$ of $y$ is given by

$$\tan \Phi_y(t) = \epsilon \tan 2\pi ft, \quad \text{with} \quad \epsilon = \frac{1 - \alpha}{1 + \alpha}. \quad (2.7)$$
On the other hand, the (unwrapped) argument of the original signal is \(\Phi_x(t) = 2\pi ft\); thus one can compute the argument distortion \(d\) as

\[
d(t) = \text{wrap}(2\pi ft) - \Phi_y(t) = \text{wrap}(2\pi ft) - \arctan(\epsilon \tan 2\pi ft),
\]

from which one can see that \(d(t) = 0\) when \(t = k/4f\), \(k = 0, 1, \ldots\); between these points, \(d\) will change its sign, which means it will oscillate at twice the tuning frequency \(f\).

If \(x(t)\) has a more complex structure (i.e., if it is a sum of sinusoids with different frequencies, amplitudes, and phases), the distortion cannot be characterized in an easy way, but it will also have a significant effect on the measured phase.

### 2.2.3 Sinusoidal quadrature filters

One alternative to Gabor filters are the sinusoidal quadrature filters (SQF’s - [Guerrero et al., 2005]) whose asymmetrical frequency response has a finite support which never intersects the negative frequencies. The frequency response of an SQF is given by:

\[
G_{\omega, h_k}(\omega) = \begin{cases} 
\frac{1}{2} [1 + \sin \left(\frac{(h_k + 2(\omega - \omega_k))\pi}{2h_k}\right)] & \text{if } \omega \in [\omega_k - h_k, \omega_k], \\
\frac{1}{2} [1 + \sin \left(\frac{(h_k + 2(\omega - \omega_k))\pi}{2h_k}\right)] & \text{if } \omega \in [\omega_k, \omega_k + h_k], \\
0 & \text{otherwise},
\end{cases}
\]

where \(\omega_k\) is the tuning frequency, \(h\) is the bandwidth, and \(h_k = \min\{h, \omega_k\}\). Figure 2.2 shows the frequency response curves for Gabor filters and SQF’s tuned at 1 Hz and 10 Hz.

The filter kernels for SQF’s can be obtained as the inverse Fourier transform (Equation 2.1) of their frequency response function. The filtered signal is then obtained by convolution with these kernels. As one can see in Figure 2.1, SQF’s do not distort the phase even for low tuning frequencies. Later, we will compare the results of the actual synchrony analysis procedure using both Gabor filters and SQF’s to determine how seriously the phase distortion (introduced by Gabor filters) affects our results.

### 2.2.4 Wavelet analysis

Wavelet analysis is equivalent to using a bank of band-pass filters, with the exception that the kernel width of the wavelet-based filters depends inversely on the tuning frequency. Lachaux et al. [1999, 2000b] use the Gabor wavelet extensively in their work. The bandwidth of the Gabor wavelets is proportional to the tuning frequency (for instance, according to [David et al., 2003a] they
Figure 2.2: Frequency response of Gabor filters (red curve) and sinusoidal quadrature filters (blue curve) tuned at 1 Hz (left graph) and 10 Hz (right graph). At relatively high tuning frequencies, both filters have a very similar response; however, at low tuning frequencies, Gabor filters show a strong response at negative frequencies, which may result in distorted phase estimates.

chose the wavelet parameters such that at a tuning frequency of 40 Hz, the bandwidth of the filter covers from 20 Hz to 60 Hz). However, since the phase obtained from the output of a filter only has a physical meaning if the signal has narrow bandwidth, it becomes necessary to pre-filter the EEG signals before using wavelets for TF decomposition.

2.3 Long-range synchrony measures

Although it is common that several areas of the brain interact together at a given time in the course of a cognitive task, synchronization measures are usually estimated only between pairs of leads. This simplifies the problem enormously, and also makes the comparison of different measures relatively simple.

In a very general sense, two simultaneous signals are synchronized in a given time window when they show a certain correlation. This correlation can be defined in different ways, depending on the physiological motivation, for example:

1. If both signals show a spike within the time window (e.g. synchrony at neuron level).
2. If their difference in phase is relatively constant (for narrow band signals).
3. If the signals are statistically inter-dependent (e.g. if their statistical correlation is high)
Thus, a synchronization measure is one that estimates the degree of correlation (according to some criterion) between the two signals. Here we describe the most common measures found in the literature.

2.3.1 Coherence

According to [Gardner, 1992], the coherence of two zero-mean random variables $X$ and $Y$ can be defined simply as the squared magnitude of their correlation coefficient $\rho_{XY}$, given by:

$$\rho_{XY} = \frac{E\{XY^*\}}{[E\{|X|^2\}E\{|Y|^2\}]^{1/2}},$$

(2.9)

where $E\{\cdot\}$ denotes the expected value. In practice, $X$ and $Y$ are usually series of $N$ sample values and the empirical degree of coherence can be estimated as follows:

$$|\rho_{XY}| = \frac{|\hat{R}_{XY}|}{[\hat{R}_{XX}\hat{R}_{YY}]^{1/2}},$$

(2.10)

with

$$\hat{R}_{XY} = \frac{1}{N} \sum_{i=1}^{N} X(i)Y^*(i).$$

(2.11)

The coherence coefficient measures the linearity of the relation between $X$ and $Y$, regardless of their difference in magnitude. It is normalized between 0 and 1, where $|\rho| = 1$ means that $Y$ can be obtained as a linear transformation of $X$.

If $X$ and $Y$ represent two time series (such as two EEG signals), it is possible to spectrally decompose the coherence by passing both $X$ and $Y$ through a tunable narrow-band bandpass filter, and then measuring the coherence of the filtered signals.

Coherence is widely used as a measure of EEG synchrony (see below), however, it is originally defined for stationary time-series. Since EEG signals are very dynamic, it is common to estimate the coherence $C_{XY}(t)$ across a short time window centered around a time of interest $t$ as follows:

$$C_{XY}(t) = \frac{|\hat{R}_{XY}(t)|}{[\hat{R}_{XX}(t)\hat{R}_{YY}(t)]^{1/2}},$$

(2.12)

with

$$\hat{R}_{XY}(t) = \frac{1}{2w+1} \sum_{i=t-w}^{t+w} X(i)Y^*(i).$$

(2.13)

The size of the time window $(2w+1)$ is typically a few hundreds of milliseconds, which corresponds to the duration of most cognitive events. By inspecting
Figure 2.3: Average coherence (taken across trials) between electrodes T5 and O2 of the Figures experiment at 8 Hz. Note the border effects caused by the large window size (which is required for reliable coherence measures); these effects must be taken into account when performing a statistical significance analysis in order to avoid biasing. Disregarding these border effects (around 150 ms on each side), one can clearly see an increase on the average level of synchronization between the electrodes under study.

or analyzing $C_{XY}(t)$, it is possible to find episodes of high coherence in an experiment. As an example, Figure 2.3 shows the average 8 Hz coherence (taken across all trials) between electrodes T5 and O2 of the Figures experiment, using a window size of 105 ms ($w = 10$ samples).

In [Bressler et al., 1993], coherence is used to study the brain integration in monkeys during a visual pattern discrimination task within a GO/NO-GO scheme: depending on the stimulus, the subjects must respond (GO condition) or withhold the response (NO-GO condition). The EEG data consisted of intracranial local field potential (LFP) recordings from 15 cortical sites. In the GO condition, Bressler et al. found episodes of significantly high coherence over a broad frequency range (from 12.5 Hz to 87.5 Hz) between distant sites. According to Bressler, “elevated coherence, rather than simply appearing because of common activation of multiple cortical areas by an imposed stimulus, manifested spatially and temporally complex patterns in the period after stimulus presentation, when the monkeys had to discriminate successfully between two visual forms and then perform correctly.”

Friston et al. [1997] suggests that dynamical correlations between two narrow band EEG signals may be related to changes in the phase-locking between the signals: using experimental magnetoencephalographic (MEG) data (only a component in the gamma band), they found that in episodes with high cross-correlation between two sites, the phase difference between both signals had a larger tendency towards zero, whereas in the low-correlation episodes, the histogram of the phase differences was much flatter. This is supported by a
theoretical analysis showing the relationship of the coherence measure and the distribution of the phase differences.

Gross et al. [2001] describe a method to study the neural interactions from the analysis of coherent sources. This method estimates the coherence between pairs of cortical sources (an inverse-problem approach is used to estimate the activity at these sources). Gross et al. studied the cortical interactions in a patient with Parkinson’s disease and found significant coherence values in the alpha band (particularly 9 to 12 Hz) between sources separated by more than 6 cm.

Another method based on the coherence of cortical sources is proposed by Hoechstetter et al. [2004]. We have already described (in Section 1.6) the inverse approach used by Hoechstetter to obtain the source signals. These signals are then passed through a bank of bandpass filters (from 4 to 50 Hz in 2 Hz steps) and coherence was estimated between pairs of sources. As an example, the method was used to analyze a task in which four different letter strings (HHHHH, SSHSS, SSSSS, and HHSSH) were presented to the subject, who was told to indicate the central letter by pressing a key with the left or right hand, respectively. Large coherence was found in the low frequency range between the left and right visual cortex regions, and also in the gamma band between the midline-visual cortex, and the left and right motor cortex areas. A widely distributed decrease in coherence was also found around the 10 Hz range.

### 2.3.2 Phase Locking Statistic

According to Varela et al. [2001], the most plausible mechanism of neural integration is the formation of dynamical links, which result in an increase of phase-locking between signals. In general, two (narrow band) signals with instantaneous phases \( \phi_1(t) \) and \( \phi_2(t) \), respectively, are phase-locked in a time window \( T \) if

\[
  n\phi_1(t) - m\phi_2(t) = \text{constant}, \quad t \in T, \quad (2.14)
\]

where \( n \) and \( m \) are integers indicating the ratios of possible frequency locking (called \( m : n \) synchrony). For simplicity, most studies concentrate on signals within the same frequency band (i.e. \( m = n = 1 \), or 1 : 1 synchrony).

Lachaux et al. [1999] introduced a method to estimate the degree of phase-locking and detect significant synchrony values between two recording sites, in a specific frequency range. The phase-locking statistic (PLS) is defined as

\[
P_{LS_{e_1,e_2}}(t) = \frac{1}{N_r} \left| \sum_{n=1}^{N_r} \exp[i(\phi_{n,e_1}(t) - \phi_{n,e_2}(t))] \right|, \quad (2.15)
\]

where \( e_1 \) and \( e_2 \) represent the two recording sites, \( N_r \) is the number of trials, and \( \phi_{n,e} \) is the instantaneous phase of the bandpass filtered signal at site \( e \) for trial \( n \). This analysis is performed for all frequency bands of interest.
Figure 2.4: PLS estimated between electrodes T5 and O2 of the Figures experiment at 8 Hz, where an increase of synchrony can be observed in the post-stimulus segment.

The PLS equals one minus the inter-trial variance of the phase differences between the two signals (see [Fisher, 1995] for a definition of circular variance). If the phase difference varies little across trials, the PLS is close to 1; otherwise, it is close to zero. Figure 2.4 shows the estimated PLS at 8 Hz for electrodes T5 and O2 from the Figures experiment. As in the previous case (with coherence), an increment of synchronization can also be observed after the stimulus onset.

In order to determine the significance of the phase-lock values, a randomized (surrogate) set of data is constructed by shuffling trials for one of the electrodes under study. This and other methods of significance estimation will be discussed in Section 2.4.

The PLS was tested with simulated data and with real human intracortical recordings (from an epileptic patient performing a visual discrimination task - see [Lachaux et al., 1999] for details). The analysis of real data focused only on one frequency band around 45 Hz, and showed significant long-range synchronizations that emerged and disappeared over time. Short-range couplings were also found; however, in this case, the PLS decreased steadily with interelectrode separation, which suggests that those couplings were spuriously introduced by volume conductor effects.

Unlike coherence, the PLS does not assume stationarity of the signals (which can rarely be validated), and does not reduce the time resolution of the analysis. Furthermore, coherence also increases with amplitude correlation, and the relative importance of amplitude and phase correlation in the coherence measure is not clear.

PLS was used by Rodriguez et al. [1999] to study gamma synchronization during an ambiguous visual discrimination task: the subjects were shown Mooney faces, which are black and white shapes that resemble a face when...
presented in upright orientation (perception condition), but are usually seen as meaningless shapes when presented upside-down (no-perception condition). Phase synchronization (within the gamma range) was studied using the PLS method, which showed differences between the perception and no-perception conditions: during perception, significant couplings between distant sites were observed that did not show in no-perception. Similarly, a global desynchronization pattern following the coupling stage was also observed only in the perception condition. Interestingly, as previously found in [Friston et al., 1997], synchronized electrodes showed a phase-lag distribution centered at zero, regardless of the interelectrode distance.

2.3.3 Single-Trial Phase Locking Statistic

One limitation of the phase-locking statistic is that it requires several trials to estimate the degree of synchrony between two signals. In some cases, it is desirable to perform a statistic with as few trials as possible, or even with single trials (for example, in experiments where a specific induced response is not necessarily time-locked to the stimulus). To overcome this limitation, the smoothed or single-trial phase-lock statistic (STPLS) was introduced [Lachaux et al., 2000b], which is given by:

\[
STPLS_{e_1,e_2}(t) = \frac{1}{2w + 1} \left| \sum_{t' = t - w}^{t + w} \exp \left[ i (\phi_{e_1}(t') - \phi_{e_2}(t')) \right] \right|.
\] (2.16)

The STPLS equals one minus the variance of the phase differences across a time window centered at time \( t \) with length \( 2w + 1 \). A STPLS of 1 means that both signals are perfectly in phase across the time window. This measure appears to be more in agreement with the synchrony criterion given by Equation 2.14, at the expense of reducing the time resolution of the analysis. Figure 2.5 shows the STPLS values for electrodes T5 and O2 of the Figures experiment, at 8 Hz. The window size is 105 ms (\( w = 10 \)). Note that the results are very similar to those obtained with coherence.

This measure has also been used by Mizuhara et al. [2005] to study neural integration during an arithmetic task. The phase-lock measure was estimated between pairs of current source density signals (see Section 1.6) and then correlated with fMRI measurements to determine which networks were involved in synchronous processes. According to the results, the fMRI showed significant responses in the parietal and temporal cortices, which were associated with phase-synchronization observed in the beta range (around 14 Hz).

Schack et al. [2005] have also used this measure to test for 1 : 1 synchrony, but also provide an extended measure to study \( m : n \) synchrony:

\[
\Gamma_{e_1,e_2}(t) = \frac{1}{2w + 1} \left| \sum_{t' = t - w}^{t + w} \exp \left[ i (m\phi_{\omega_n,e_1}(t') - n\phi_{\omega_m,e_2}(t')) \right] \right|,
\] (2.17)
Figure 2.5: Average STPLS (estimated across all trials) between electrodes T5 and O2 of the Figures experiment at 8 Hz, where an increase of average synchrony can be observed in the post-stimulus segment (disregarding the border effects). These results are very similar to those obtained with coherence.

where \( \phi_{\omega_n,e} \) is the instantaneous phase of the signal at site \( e \), filtered around frequency \( \omega_n \). There must be a harmonic relation between the frequencies \( \omega_n \) and \( \omega_m \).

This measure was used to study a working memory task where a set of 1 to 4 digits were presented to each subject, followed by an ‘x’ symbol indicating the beginning of the retention interval (see [Schack et al., 2005]). After the retention interval, another digit (probe) was presented and the subjects were instructed to respond with a keypress whether the probe was or was not in the set. Significant 1 : 1 synchrony was found in the theta (6 Hz) and upper alpha (12 Hz) bands between distant sites (e.g. right-frontal vs. left-temporal). Inter-frequency \((m : n)\) synchronization was also found between the sites F7 and O2 of the 10/20 system, corresponding to theta/upper-alpha couplings \((1 : 2\) ratio).

2.3.4 Generalized Synchronization

According to David et al. [2004], two dynamical systems \( X \) and \( Y \) show generalized synchronization (GS) when “the state of the response system \( Y \) is a function of the state of the driving system \( X \) : \( Y = F(X) \)”. If \( F \) is continuous, then close points on the attractor of \( X \) should correspond to close points on the attractor of \( Y \).

The method described in [Quian-Quiroga et al., 2002] and [David et al., 2004] to estimate the GS relies upon vectors \( \vec{X}_n = [x_n, x_{n+\tau}, \ldots, x_{n+(m-1)\tau}] \) and \( \vec{Y}_n = [y_n, y_{n+\tau}, \ldots, y_{n+(m-1)\tau}] \), which are embeddings of the signal \( x \) in an \( m \)-dimensional space with a delay time \( \tau \). If \( p_{ni} \) and \( q_{ni}, i = 1, \ldots, k, \) denote the indices of the \( k \) nearest neighbors of \( \vec{X}_n \) and \( \vec{Y}_n \), respectively, then one can define a measure of closeness \( S^n_k(X|Y) \) between the true neighbors \( \{\vec{X}_{p_{ni}}\} \) of
\( \vec{X}_n \) and its mutual neighbors \( \{ \vec{X}_{qni} \} \). Quian-Quiroga proposes the following measure:

\[
S_k^X(X|Y) = \frac{\langle |\vec{X}_n - \vec{X}_{pni}|^2 \rangle_i}{\langle |\vec{X}_n - \vec{X}_{qni}|^2 \rangle_i},
\]

(2.18)

where \( \langle \cdot \rangle_i \) indicates the average value across \( i = 1, \ldots, k \).

Equation 2.18 measures the ratio of the mean square distance between \( \vec{X}_n \) and its \( k \) nearest neighbors, and the mean square distance between \( \vec{X}_n \) and its mutual neighbors (with respect to \( \vec{Y}_n \)). By construction it is bounded between 0 and 1, where 1 means that the cloud of the \( k \) nearest neighbors of \( \vec{Y}_n \) maps exactly to the cloud of \( k \) nearest neighbors of \( \vec{X}_n \). Finally, synchronization measure is the average of \( S_k^X(X|Y) \) over a time window \( t - T \leq n \leq t + T \).

According to Quian-Quiroga, GS should be very sensitive to weak correlations (which may or may not be useful for specific applications) - this is confirmed by experiments performed by David et al. On the other hand, GS is also computationally demanding and greatly reduces the time resolution of the synchrony analysis, which may be a problem when studying the gamma band, where the synchronization periods may be relatively short. Another disadvantage of GS is that it relies on many parameters: \( \tau, m, k \), and \( T \), which control the tradeoff between the robustness, time-resolution, and computational load of the method. Unfortunately, neither David or Quian-Quiroga provide a comparison of the behavior of GS for different parameter values.

### 2.3.5 Mutual Information

Mutual information (MI) is a statistical measure which estimates, for two random variables \( X \) and \( Y \), the amount of information in \( X \) given that \( Y \) is known, and vice versa. MI is based on the entropy \( H \) of a discrete random variable \( X \in \{ x_1, \ldots, x_M \} \), which is given by

\[
H(X) = - \sum_{i=1}^{M} p_i \ln p_i,
\]

(2.19)

where \( p_i = P(X = x_i) \). If \( X \) is continuous with a finite support, one can simply partition the support in \( M \) bins and estimate the \( p_i \)'s by counting. For a bi-variate distribution, the joint entropy is defined as

\[
H(X,Y) = - \sum_{i,j} p_{ij} \ln p_{ij},
\]

(2.20)

where \( p_{ij} = P(X = X_i, Y = Y_j) \). If \( X \) and \( Y \) are independent, we have that \( H(X,Y) = H(X) + H(Y) \).

The mutual information between \( X \) and \( Y \) is defined as

\[
MI(X,Y) = H(X) + H(Y) - H(X,Y).
\]

(2.21)
If $X$ and $Y$ are independent, then $MI(X, Y) = 0$; otherwise it will take positive values up to $H(X)$. MI basically measures the similarity between the histograms of $X$ and $Y$ (estimated, in our case, across a time window), thus it does not take into account any possible time lag between both signals. This makes it inadequate as a phase-locking measure. It is possible, however, to apply time-embedding techniques (for example, taking as sample data the vectors $\vec{X_n}$ used in GS) to obtain a lag-sensitive MI measure [Quian-Quiroga et al., 2002].

MI presents the same problems that GS: it is computationally expensive (compared to phase-locking measures such as coherence or PLS), and if time-embedding techniques are used, it will be at the expense of high time resolution. For these reasons, MI and GS are better suited for the analysis of stationary couplings and/or low frequency synchrony.

### 2.4 Statistical significance analysis

Whether one is studying evoked potentials, power changes, or synchrony between pairs of electrodes, it is likely that one will obtain a set of time series $\{X_k(t)\}$, each of which represents the computed measure for a specific combination of trial number, frequency band, electrode, or electrode pair (represented by $k$). For example, in the case of evoked potentials, $k$ may represent an electrode index, and $X_k$ would be the average signal at that electrode, whereas in the case of synchrony, $k$ may represent a 4-tuple of the form $< j, \omega, e_1, e_2 >$ for trial $j$, frequency $\omega$, and electrode pair $< e_1, e_2 >$.

Once a specific measure $X(t)$ is computed, one would like to determine, for each $t$, if $X(t)$ represents a significant deviation from what would be considered as a null condition (for example, if one is studying event-related activity, one can consider the pre-stimulus segment as the null condition). To do this, one can estimate the distribution of $X$ during the null condition, and then perform a statistical test to determine if $X(t)$, for any $t$, comes from the null distribution.

#### 2.4.1 Choosing the null population

The null population consists of those values of $X(t)$ which are observed during the null condition. For example, if one wants to study task-related activity, one can assume that everything that happens before the stimulus onset is part of the null condition, and significant changes in $X$ with respect of this condition may be related with the task. In this case, one choice for the null population is simply those values of $X$ in the pre-stimulus (i.e., $X(1), \ldots, X(T_s)$ - although in practice, possible border effects due to filtering or averaging should be taken into account). For example, this is the method used by [Marroquin et al., 2004] to estimate the significance of power changes (i.e., induced power) during a cognitive task. A similar technique was used by [Mizuhara et al., 2005].

It is also common to determine if the $X$ values are significantly different from what can be expected by chance. In this case, one can build the null population
by taking the original data and manipulating it in order to obtain a new set of surrogate data which does not have the property of interest (i.e., the property which is being measured). In particular, when one is measuring synchrony between two signals, one would like to construct the null population so that the two surrogate signals are certainly unsynchronized. This can be achieved, for example, by shuffling the samples in each signal, which would destroy any temporal structure the signals may have. One can then compare the measured synchrony against the distribution of the surrogate data, and decide if there is significantly more synchrony than what can be expected by chance. Lachaux et al. [1999; 2000] use surrogate data to estimate the significance of their PLS and STPLS measures: the surrogate synchrony values, for two electrode sites $e_1$ and $e_2$, is obtained by shuffling the trials for $e_2$.

Once the null population is obtained, one can estimate its distribution in various ways, or use statistical tools, such as ANOVA, to compare the $X$ values against the null condition. We will focus on estimating the null distribution and estimating the $p$-values for each $X(t)$.

### 2.4.2 Significance estimation: normal distribution

One of the most common methods used to determine the significance of the synchrony changes consists in assuming that the null distribution is normal with mean $\bar{X}_{null}$ and variance $\sigma_{null}^2$, and computing the $Z$ variables given by:

$$Z(t) = \frac{X(t) - \bar{X}_{null}}{\sigma_{null}}. \quad (2.22)$$

In practice, if the size $n$ of the null population is sufficiently large (e.g., $n > 30$), one can replace $\bar{X}_{null}$ and $\sigma_{null}^2$ with the sample average and sample variance of the null population, respectively.

Under the null condition, $Z$ is normally distributed with mean 0 and variance 1. Therefore, given the significance level $\alpha$, the quantile $z_{(1-\alpha)}$ represents the threshold that the $Z$ values must cross to be considered significantly different. For example, if $Z(t) > z_{(1-\alpha)}$, then $Z(t)$ is significatively greater than in the null condition. Common values for $\alpha$ are 0.05 and 0.01, corresponding to thresholds $z_{0.05} = 1.64$, and $z_{0.01} = 2.33$.

It is also possible to test the average $\bar{X}(t)$ of $X$ across a time window against the null condition. For example, if the time window goes from $t - w$ to $t + w$, $\bar{X}(t)$ would be given by

$$\bar{X}(t) = \frac{1}{2w + 1} \sum_{t'-t-w}^{t+w} X(t'), \quad (2.23)$$

which is normally distributed with mean $\bar{X}_{null}$ and variance $\sigma_{null}^2/(2w + 1)$. Therefore, the $Z$ variables would be given, in this case, by

$$Z(t) = \frac{\bar{X}(t) - \bar{X}_{null}}{\sigma_{null}/\sqrt{2w + 1}}. \quad (2.24)$$
2.4.3 Significance estimation: kernel density estimation

It may not be correct to assume normality for the $X$-values, especially if the number of trials is relatively small. In this case, the null distribution $p_X$ can be estimated from the null population $\{x_i\}$ as the sum of superimposed kernel functions $k_h$ centered at each $x_i$. In other words,

$$p_X(x) = \frac{1}{Z} \sum_{i=1}^{n} k_h(x - x_i),$$

where $n$ is the sample size, and $Z$ is a normalization constant chosen so that $p_X$ integrates to 1. The parameter $h$ specifies the width of the kernel and it determines the smoothness of $p_X$. Adequate choice of $h$ depends on the sample data; however, for automated applications, a common rule of thumb for a Gaussian kernel (which we use) is provided by [Silverman, 1986]:

$$h_{\text{opt}} = 1.06 \sigma n^{-1/5},$$

where $\sigma$ is the standard deviation of the null population (usually estimated from the sample).

The p-value of a given $x^*$ is then defined as $p_X(x > x^*) = 1 - p_X(x < x^*) = 1 - P_X(x^*)$ where $P_X(x)$ is the cumulative density function (cdf) corresponding to $p_X$, which can be defined as:

$$P_X(x^*) = \int_{-\infty}^{x^*} p(x) dx. \quad (2.25)$$

If the p-value for a given $x^*$ is lower than the significance level $\alpha$, then $x^*$ is considered to be significative higher with respect to the null distribution. A similar test can be performed to determine if $x^*$ is significatively lower.

In practice, one can simply obtain $p_X$ by convolving the histogram (with equally spaced bins) of the null population with the gaussian kernel, and normalizing the resulting histogram so that it integrates to 1. One can then approximate the cdf using numerical integration methods. For example, given a histogram $(h, b)$ where $h_n$ represents the frequency (count) of samples which fall in the range $[b_n, b_{n+1})$ (i.e., the $n$-th bin), the cdf $P_X(x^*)$ can be approximated using rectangular integration (where each bin defines a rectangle whose height is $h_n$) as follows:

1. Compute $h'$ as a normalized version of $h$; that is, $h'_n = h_n / \sum_i h_i$. $h_n$ represents the probability of $x$ falling into bin $n$.

2. Find $j$ such that $b_j \leq x^* < b_{j+1}$.

3. Estimate the cdf as $P_X(x^*) = h'_j (x^* - b_j) / (b_{j+1} - b_j) + \sum_{i<j} h'_i$. 

30
2.5 Visualization techniques

Synchrony values are usually estimated for various electrode pairs, at different frequencies, and for any point in the timeline. In other words, the data resulting from a synchrony analysis is 4-dimensional, which represents a visualization problem. Because of this, most works resort to averaging across time or frequency, or focus only on specific regions of the 4-dimensional synchrony space, in order to display their results.

The simplest type of display is a typical XY graph of a single-variable function. For example, one may take a fixed electrode pair and frequency, and plot the corresponding coherence time-series, or their significance values. However, this type of graph does not reveal anything about the spatial or spectral distribution of synchrony values. Other works use a matrix display of a two-variable function, in which the color assigned to each point represents the synchrony value. Typically, one fixes the electrode pair and displays a time-frequency map (an example is presented in Figure 2.6a). Many works use this type of display (Varela et al., 2001), Hoechstetter et al., 2004, Schack et al., 2005), but one may also choose a fixed time-frequency window and display the synchrony values for each electrode pair (Lachaux et al., 1999).

Another common type of display (shown in Figure 2.6b), for a fixed time-frequency region, consists on a head diagram where each electrode site is represented by a dot. Significant synchrony between two sites is represented by a straight line connecting the corresponding dots (Rodriguez et al., 1999, Varela et al., 2001, Mizuhara et al., 2005, Schack et al., 2005). The problem with this display is that, when there is a considerable amount of synchronous activity, too many lines overlap and it becomes difficult to grasp the spatial distribution of the synchrony values. An alternative consists on a multitoposcopic display, in which at each recording site , one shows a head diagram with the distribution of synchrony between and every other site (Figure 2.6c).

There may be difficulties when one wants to plot a multi-toposcope for high electrode density recordings: only a certain number of toposcopes (head diagrams) can be presented within the bigger head before the graph becomes too cluttered. In Chapter 3 we will offer a few suggestions to overcome this problem.

Marroquin et al. introduced in 2004 a method for the study of EEG-based psychophysiological experiments, which analyzes power changes in EEG and presents the results in a detailed time-frequency-topography (TFT) display [Marroquin et al., 2004]. The methodology basically consists of four parts:

1. A time-frequency decomposition using Gabor filters, from which the log-power signals are obtained.

2. Estimation of a statistic , which measures the mean deviation of the log-power with respect to the pre-stimulus average, for each frequency and electrode .
Figure 2.6: Common displays used for synchrony data: (a) Time-Frequency map of significance indexes of synchrony changes with respect to a null condition (in this case, the pre-stimulus segment) during a figure classification task ([Marroquin et al., 2004]). The data corresponds to the electrode pair Fp1-O2. Red regions correspond to significant increases in synchrony, whereas green regions indicate significant decreases. (b) Synchrony patterns observed at specific time-frequency points during the figure classification task. Only significant increases in synchrony with respect to the pre-stimulus average are shown. (c) Multitopososcopic representation of the corresponding synchrony patterns shown in (b). Both increases (red) and decreases (green) are shown.
3. Estimation of a significance index $I_{\omega,e}(t)$ based on the p-value of $Y_{\omega,e}(t)$ with respect to the distribution of $Y_{\omega,e}$ in the pre-stimulus.

4. Plot the significance indices as follows: for each time $t$ and frequency $\omega$, draw a small head diagram with the distribution of the significance values $I_{\omega,e}(t)$ for all electrodes (see Figure 2.7).

Additionally, one may perform a segmentation of the time-frequency (TF) plane in regions with homogeneous activation patterns (that is, patterns of simultaneous increases and decreases of power which are significant). For this, Marroquin et al. propose a Bayesian approach based on a Markov Random Field (MRF) model of a class field $C_{\omega,e}(t)$, which indicates if the $Y$-values are significantly higher (class $C = 1$), lower ($C = -1$), or equal ($C = 0$), with respect to the pre-stimulus distribution. A regularization constraint is included in the model in order to favor solutions with relative large regions with homogeneous activation patterns.

While this procedure is designed for amplitude analysis, some of the techniques it involves can also be used for synchrony analysis; particularly the display techniques, since synchrony data is highly dimensional.
Chapter 3

Phase synchrony analysis

3.1 Procedure overview

In this chapter, we describe in detail the methodology we follow for the analysis of EEG synchrony patterns. Our procedure consists of the following steps:

1. Run the EEG signals through a bank of bandpass quadrature filters and extract phase information (TF phase analysis).
2. Calculate a synchrony measure from the filtered signals.
3. Estimate the likelihoods and prior distributions for the MRF model using the synchronization values.
4. Use Bayesian estimation to find significant synchronization patterns that are persistent.
5. Display synchronization patterns as multitoposcopic graphs and time-frequency-topography (TFT) maps.

We illustrate our procedure using data from a figure classification experiment [Harmony et al., 2001] where white-line figures on a black background were presented to each subject. The subjects were instructed to press a button if the figure corresponded to an animal whose name started with a consonant, and another button if the figure did not correspond to an animal and the name of the figure started with a consonant. If the name started with a vowel, the subject was instructed not to respond. The subjects were 18 normal children (8 to 10 years old, 9 females), all right handed with normal neurological examination. EEG was recorded with reference to linked ears from Fp1, Fp2, F3, F4, C3, C4, P3, P4, O1, O2, F7, F8, T3, T4, T5, T6, Fz, Cz, Pz, and Oz of the 10/20 system. EOG was recorded from a supraorbital electrode and from an electrode on the external canthus of the right eye. The amplifier bandwidth was set between 0.5 and 30 Hz. EEG was sampled every 5ms using a MEDICID 3E system and
stored on a hard disk for further analysis. Sampling was done every 5 ms during a time segment from 1280 ms before the stimulus to 1500 ms after its onset. Each trial was visually edited and only those corresponding to correct responses and with no artifacts were analyzed. Subjects were seated in a comfortable chair in front of the videomonitor. Stimuli were delivered by a MINDTRACER system synchronized to the MEDICID 3E acquisition system.

The recorded potentials will be denoted by $V_{j,e,t} = V_{j,e}(t)$, where $j = 1, \ldots, N_r$ is the trial number, $e = 1, \ldots, N_e$ represents the electrode site, and $t = 0, \ldots, N_t - 1$ is the time index (in samples), with $t = 0$ indicating the beginning of each recording. The time of stimulus onset (pre-stimulus length) will be denoted by $T_s$. In particular, for the Figures experiment we have $N_r = 1290$, $N_e = 20$, $N_t = 512$, and $T_s = 200$.

The procedure has also been tested with other various experiments: a word categorization experiment ([Harmony et al., 2001]) similar to the Figures experiment (with words presented instead of figures), a working-memory task based on the Sternberg paradigm ([Harmony et al., 2004]), and a Go/NoGo experiment to study the inhibition of the motor response ([Harmony et al., 2006]). Details and results for these experiments can be found in Chapter 6.

3.2 Instantaneous phase extraction

In order to extract phase data from the EEG signals, one can perform a time-frequency decomposition using a bank of quadrature filters. We have already studied two possible choices: Gabor filters and Sinusoidal Quadrature filters (SQF's). We have chosen the Sinusoidal filters to avoid any distortion that may be introduced by Gabor filters tuned at low frequencies. Later, in Section 3.6, we will compare the results of the analysis described below, using both types of filters, and determine if there is an actual advantage of SQF’s over Gabor filters.

In general, one can obtain a time-frequency decomposition of the EEG signals by convolving each $V_{j,e}$ with a bank of bandpass filters whose kernels are $K_\omega$, where $\omega$ is the tuning frequency of the filter. In other words, the filtered signal $F_{\omega,j,e}(t)$ is given by:

$$F_{\omega,j,e} = V_{j,e} * K_\omega,$$

We have chosen a bank of filters tuned at each Hertz, from 1 Hz to $N_f = 40$ Hz, with a fixed bandwidth for all the filters, equivalent to a standard deviation $\sigma_\omega \approx 1$ Hz ($\sigma_t = 15$ samples) for the Gaussian response of Gabor kernels. This results in a bandwidth of 1.76 Hz within 3 dB of attenuation. The width of Sinusoidal filters was chosen to have a similar bandwidth than Gabor filters ($h = 5\sigma_\omega$, approximately).
Finally, one can extract, from the filtered signals, the instantaneous amplitude \( \hat{A} \) and argument \( \hat{\Phi} \), which are given by

\[
\hat{A}_{j,\omega,e}(t) = \sqrt{\Re^2(F_{j,\omega,e}(t)) + \Im^2(F_{j,\omega,e}(t))},
\]

(3.2)

\[
\hat{\Phi}_{j,\omega,e}(t) = \text{atan2}(\Im(F_{j,\omega,e}(t)), \Re(F_{j,\omega,e}(t))],
\]

(3.3)

with \( \hat{\Phi}_{j,\omega,e}(t) \in [-\pi, \pi) \).

The output of a sufficiently narrow bandpass filter can be approximated by a sinusoidal whose frequency is the tuning frequency of the filter \( \omega \), and whose argument can be expressed as \( \Phi(t) = \omega t + \phi(t) \), where \( \phi(t) = \Phi(t) - \omega t \) is the phase of the signal. While it is sometimes necessary to work with the phase instead of the argument of the signals, most phase-synchrony measures are based on the phase difference \( \phi_1 - \phi_2 \) between two signals, thus one can simply use \( \Phi_1 - \Phi_2 \) since the \( \omega t \) terms cancel each other.

### 3.3 Instantaneous phase-lock measures

Here we propose two measures for in-phase synchrony (i.e., synchrony with a phase difference of zero). As a convention, we will normalize all synchrony measures between 0 and 1, so that 1 means perfect synchrony. All measures presented in Section 2.3, with the exception of Mutual Information, follow this convention.

#### 3.3.1 Motivation

In 2003, David and Friston introduced a neural mass EEG model which simulates the activity of populations of neurons, under the assumption that the state of the population can be approximated with very few state variables [David and Friston, 2003b]. According to this model, the EEG/MEG signals from two distant cortical areas will show a phase difference of 0 (in-phase) or \( \pi \) (anti-phase) when there is a bidirectional coupling between the two areas. According to David and Friston [2003a], and Varela et al. [2001], the proportion of reciprocal connections in the brain is very high, suggesting that EEG synchrony may be predominantly in-phase and anti-phase synchrony. Moreover, there is experimental evidence supporting the idea that the distribution of the phase differences concentrates around zero during episodes of high synchrony (see, for example, [Friston et al., 1997; Rodriguez et al., 1999]). Therefore, it makes sense to consider as synchrony criterion the following: two signals with instantaneous phases \( \phi_1(t) \) and \( \phi_2(t) \) are in synchrony when \( \phi_1(t) \approx \phi_2(t) \) for all \( t \) in a given time interval.
In Chapter 4 we will show that synchrony measures which are estimated across a large time window, such as coherence and STPLS, may be contaminated by the dispersion of the phase at each electrode, which is not necessarily related to the degree of synchrony between the pair of signals under study. For this reason, we favor synchrony measures which are “instantaneous”, in the sense that they are estimated from instantaneous values of the filtered signals (disregarding that the filtering process itself reduces the time resolution). The rest of this section deals with two synchrony measures specifically designed for instantaneous in-phase couplings.

3.3.2 Mean Phase Difference

A straightforward instantaneous in-phase synchrony measure is simply the magnitude of the phase difference $|\phi_1(t) - \phi_2(t)|$, which can be wrapped between $-\pi$ and $\pi$. This makes it easy to obtain a normalized measure based on the phase difference:

$$
\mu_{j,\omega,e_1,e_2}(t) = 1 - \frac{1}{\pi} |\text{wrap}(\phi_{j,\omega,e_1}(t) - \phi_{j,\omega,e_2}(t))|,
$$

where $\text{wrap}(\phi)$ returns the angle $\phi$ wrapped to the interval $[-\pi, \pi)$.

Figure 3.1 shows the synchronization values between sites T5 and O2, obtained with MPD at 8 Hz. As in the previous examples (for coherence, PLS, and STPLS), an increase of synchronization between these two electrodes is shown in the post-stimulus segment. Moreover, these results are very similar to those obtained with the PLS measure, even though PLS measures synchrony with a consistent phase-lag (across trials) which is not necessarily zero. This will be further discussed in Chapter 4.
3.3.3 Cumulative Probability of Phase Difference

An alternative to the MPD measure consists on estimating the probability of the phase difference being smaller (in absolute value) than some $\epsilon > 0$. This probability can be estimated across all trials for each $t, \omega, e_1$, and $e_2$ as follows:

$$\mu_{\omega, e_1, e_2}(t) = \frac{1}{N_r} \sum_{j=1}^{N_r} I(|\text{wrap}(\phi_{j, \omega, e_1}(t) - \phi_{j, \omega, e_2}(t))| < \epsilon).$$  \hfill (3.6)

where $I(P)$ equals 1 if $P$ is true, and zero otherwise.

This measure allows a quantification of the degree of synchrony (by means of $\epsilon$), regardless of any further transformation applied to the measure (such as the significance analysis we perform). We have used $\epsilon = \pi/5$ rad for our tests (equivalent to 10 ms at 10 Hz), although one can also use a frequency-dependent value of $\epsilon$ given in time units.

CPPD synchronization between electrodes T5 and O2 from the Figures experiment is shown in Figure 3.2 (for $\omega = 8$ Hz).

One can modify this measure to detect anti-phase synchrony (couplings which show a phase difference of $\pi$). A simple example is given by:

$$\mu_{\omega, e_1, e_2}(t) = \frac{1}{N_r} \sum_{j=1}^{N_r} I(|\text{wrap}(\phi_{j, \omega, e_1}(t) - \phi_{j, \omega, e_2}(t) - \pi)| < \epsilon).$$  \hfill (3.7)

However, one must be careful not to confuse a decrease in synchrony with anti-phase couplings. This may be done, for example, by using small values for $\epsilon$.

Some preliminary results with anti-phase measures will be presented in Section 6.4.2.
3.4 Significance estimation

Since we are interested in event-related activity, we must determine how significant are the changes of synchrony with respect to the pre-stimulus segment. Following the procedure introduced by Marroquin to estimate the relative changes of EEG amplitudes [Marroquin et al., 2004], we subtract the average synchrony in the pre-stimulus segment in order to obtain the relative synchrony $X_{j,\omega,e_1,e_2}$:

$$X_{j,\omega,e_1,e_2}(t) = \mu_{j,\omega,e_1,e_2}(t) - \frac{1}{T_s} \sum_{t'=1}^{T_s} \mu_{j,\omega,e_1,e_2}(t'),$$  \hspace{1cm} (3.8)

where $T_s$ is the length of the pre-stimulus segment. Finally, we take the mean relative synchrony across all trials:

$$Y_{\omega,e_1,e_2}(t) = \frac{1}{N_r} \sum_{j=1}^{N_r} X_{j,\omega,e_1,e_2}(t).$$  \hspace{1cm} (3.9)

In the case of the CPPD measure (and also for Lachaux’s PLS), $\mu_{\omega,e_1,e_2}(t)$ itself represents an average across trials, thus $Y$ may be defined simply as

$$Y_{\omega,e_1,e_2}(t) = X_{\omega,e_1,e_2}(t) = \mu_{\omega,e_1,e_2}(t) - \frac{1}{T_s} \sum_{t'=1}^{T_s} \mu_{\omega,e_1,e_2}(t').$$  \hspace{1cm} (3.10)

Note that the mean of $Y_{\omega,e_1,e_2}$ in the pre-stimulus is zero, for every $\omega$, $e_1$, and $e_2$. Positive values of $Y$ represent synchrony increases with respect to the baseline (pre-stimulus), and vice-versa.

To facilitate interpretation, we would like to classify each value of the mean relative synchrony in one of three classes:

- Significantly higher than the baseline (class $c = 1$). This corresponds to values of $Y$ which are significantly greater than zero.
- Significantly lower than the baseline (class $c = -1$), corresponding to $Y$-values significantly lower than zero.
- No significant change ($c = 0$).

Here we will discuss two different ways to perform this classification.

3.4.1 Thresholded p-values from estimated distributions

A typical approach to determine the significance of the $Y$-values is the thresholded p-value technique described in Section 2.4.3. In this case, the null population consists of the $Y$ values in the pre-stimulus segment (disregarding samples
for \( t < 2\sigma_t \), which may be influenced by border effects). With these samples one can use kernel density estimation to construct the distribution \( p_0 \) of \( Y \) in the pre-stimulus, which can also be considered the distribution of \( Y \) given that \( c = 0 \) (i.e., \( P(Y \mid c = 0) \)). Then one can estimate a significance index \( S \) for each \( Y \), based on the \( p \)-value, as follows:

\[
S_{\omega,e_1,e_2}(t) = \begin{cases} 
1 - p_0(Y > Y_{\omega,e_1,e_2}(t) \mid Y > 0) & \text{for } Y_{\omega,e_1,e_2}(t) > 0, \\
- (1 - p_0(Y < Y_{\omega,e_1,e_2}(t) \mid Y < 0)) & \text{for } Y_{\omega,e_1,e_2}(t) < 0.
\end{cases}
\] (3.11)

The conditional probabilities in Equation 3.11 can be computed from the cumulative density function \( P_0(Y^*) = p_0(Y < Y^*) \) (see Section 2.4.3) as follows:

\[
p_0(Y > Y_{\omega,e_1,e_2}(t) \mid Y > 0) = \frac{1 - P_0(Y_{\omega,e_1,e_2})}{1 - P_0(0)}, \tag{3.12}
\]

\[
p_0(Y < Y_{\omega,e_1,e_2}(t) \mid Y < 0) = \frac{P_0(Y_{\omega,e_1,e_2})}{P_0(0)}. \tag{3.13}
\]

The significance indices represent the direction of the change of synchrony with respect to the baseline (positive means increment, whereas negative means decrement), and the degree of significance of these changes. They can be used directly to obtain the class label field \( c \) by simply comparing them with a significance threshold \((1 - \alpha)\):

\[
c_{\omega,e_1,e_2}(t) = \begin{cases} 
1 & \text{if } S_{\omega,e_1,e_2}(t) > (1 - \alpha), \\
-1 & \text{if } S_{\omega,e_1,e_2}(t) < -(1 - \alpha), \\
0 & \text{otherwise.}
\end{cases}
\] (3.14)

### 3.4.2 Bayesian classification of significant changes in synchrony

A powerful technique for classification problems consists of Bayesian estimation with a prior Markov Random Field (MRF) model [Marroquin, 1987]. With this method, one can model the class field \( c_{\omega,e_1,e_2,t} \) as a random field with a prior Gibbs distribution of the form

\[
P_{MRF}(c) = \frac{1}{Z} \exp \left[ -\lambda \sum_C V_C(c) \right], \tag{3.15}
\]

where \( Z \) is a normalizing constant and \( V_C \) is a potential function that depends only on the values of the sites belonging to the clique \( C \) (see [Marroquin, 1987, 2001] for more details). For a classification problem, where \( c \) is discrete, a
popular model is the Ising model, which enforces $c$ to be piece-wise constant. If we consider a first-order neighborhood system, whose cliques are single sites and nearest-neighbor pairs of sites, the Ising potentials are given by:

$$V_{r,s}(c) = \begin{cases} 
-1, & \text{if } c_r = c_s \\
1, & \text{if } c_r \neq c_s
\end{cases} \tag{3.16}$$

where $r$ and $s$ are nearest neighbors. In our case, $r$ and $s$ are generally 4-tuples of the form $r = (\omega, e_1, e_2, t)$; however, at this point we are only interested in modeling persistence in time, although persistence across frequencies and spatial regularization could be taken into account to attempt an automated segmentation process.

The label field $c$ is assumed to be Markovian (with a first-order Ising model) with distribution

$$P_c(c) = \frac{1}{Z} \exp \left[ -\lambda \sum_{<r,s>} V_{r,s}(c) + \sum_r \log \alpha_{c_r} \right],$$

where $\alpha_k$ represents the global prior probability $\alpha_k = P(c_r = k)$. Besides the prior distribution for $c$, we also need the likelihood $P(Y \mid c)$, which can be written as

$$P(Y \mid c) = \prod_r P(Y_r \mid c_r) = \exp \left[ \sum_r \log P(Y_r \mid c_r) \right]. \tag{3.17}$$

Using Bayes rule, the posterior distribution of $c$ given $Y$ can be calculated as

$$P(c \mid Y) = \frac{1}{Z'} \exp \left[ \sum_r \log h_r(c_r) - \lambda \sum_{<r,s>} V_{r,s}(c) \right], \tag{3.18}$$

where $Z'$ is a normalization constant and $h_r(k) = P(Y_r \mid c_r = k)\alpha_k$.

Given the estimator $\hat{c}$ and the true (unknown) field $c$, one can define a cost function $C(c, \hat{c})$ and find the optimal $\hat{c}$ by minimizing the expected value of $C(c, \hat{c})$. Since $c$ is discrete, a suitable cost function given by Marroquin is

$$C(c, \hat{c}) = \sum_r [1 - \delta(c_r - \hat{c}_r)], \tag{3.19}$$

and its expected value would be given by

$$E[C(c, \hat{c})] = \sum_c C(c, \hat{c}) P(c \mid Y)$$

$$= \sum_c \sum_r [1 - \delta(c_r - c'_r)] P(c \mid Y)$$

$$= K - \sum_{r} \sum_{c_r = c'_r} P(c_r \mid Y). \tag{3.20}$$
The posterior marginal distribution for site \( r \) is defined as:

\[
\pi_r(k) = \sum_{c: c_r = k} P(c \mid Y). \tag{3.21}
\]

Thus the optimal estimator \( \hat{c} \) (which minimizes \( E[C(c, \hat{c})] \)) can be found by maximizing \( \pi_r(\hat{c}_r) \) for each \( r \). This estimator is known as the Maximizer of Posterior Marginals (MPM) estimator and is usually approximated using stochastic Markov-chain methods such as Metropolis or the Gibbs sampler. These algorithms, however, are computationally expensive and require an unknown number of iterations, which makes them less than adequate for our multidimensional data set. A better solution consists on approximating the posterior marginal distributions with the empirical marginals \( \text{\(p\)}_r(k) \) [Marroquin et al., 2001]. It can be shown that the \( \text{\(p\)}_r \) vectors form a MRF with the same neighborhood system as \( c \), thus the distribution of \( \text{\(p\)} \) is given by

\[
P_p(p) = \frac{1}{Z_p} e^{-U(p)}, \tag{3.22}
\]

where, for a first-order neighborhood system, \( U(p) \) can be written as

\[
U(p) = \sum_r |p_r - \hat{p}_r|^2 + \lambda' \sum_{<r,s>} |p_r - p_s|^2, \tag{3.23}
\]

with \( \hat{p}_r = h_r(k)/\sum_{k'} h_r(k') \).

If \( c \) follows the Ising model, it can be shown that the field \( p \) can be modeled as a set of decoupled membrane models \( p(k) = \{p_r(k), \forall r \} \) for \( k = -1, 0, 1 \). Therefore, the optimal \( p \) is obtained by minimizing, for each layer \( k \), the energy function \( U_k(p) \) given by

\[
U_k(p) = \sum_r (p_r(k) - \hat{p}_r(k))^2 + \lambda' \sum_{<r,s>} (p_r(k) - p_s(k))^2. \tag{3.24}
\]

Since each \( p(k) \) is continuous, \( U_k(p) \) can be minimized by solving the linear system obtained from equating the partial derivatives of \( U_k \) with respect to \( p_r(k) \) to zero. However, the optimal \( p(k) \) is a smoothed version of \( \hat{p}(k) \), thus one can achieve a similar result by simply low-pass filtering each \( \hat{p}(k) \) with a Gaussian kernel (for more details see [Marroquin et al., 1997]). Once we have \( p \) we can obtain the approximated MPM estimator as

\[
c_r = \text{argmax}_k \{p_r(k)\}. \tag{3.25}
\]

### 3.4.3 Estimation of prior distributions and likelihoods

For now, we are only interested in modeling persistence in time, thus we may estimate the time series \( c_{\omega, e_1, e_2}(t) = c_{\omega, e_1, e_2, t} \) in a decoupled manner for each
\( \omega, e_1, \) and \( e_2. \) To simplify things in this section, we will consider a fixed frequency \( \omega \) and electrode pair \((e_1, e_2)\) and only keep the time subindex.

In order to calculate \( h_t(k) \) (and thus \( p_t(k) \)) we need the prior probabilities \( \alpha_k = P(c = k) \) and likelihoods \( P(Y_t \mid c = k) \). These can be estimated from the data if we consider that the complete distribution \( P(Y_t) \) can be expressed as:

\[
P_Y(Y_t) = \sum_{k=-1}^{1} \alpha_k P(Y_t \mid c = k),
\] (3.26)

and also consider the following assumptions:

- \( P(Y_t \mid c = 0) \) may be estimated empirically from the pre-stimulus data.
- \( P(Y_t \mid c = 1) = 0 \) for \( Y \leq 0 \).
- \( P(Y_t \mid c = -1) = 0 \) for \( Y \geq 0 \).

With these assumptions, we can obtain \( P(c = 0) \) from Equation (3.26) as follows:

\[
\alpha_0 = \frac{P_Y(0)}{P(0 \mid c = 0)}
\] (3.27)

and also

\[
h_t(0) = \alpha_0 P(Y_t \mid c = 0)
\] (3.28)

\[
h_t(1) = \begin{cases} P_Y(Y_t) - h_t(0), & Y_t > 0 \\ 0, & Y_t \leq 0 \end{cases}
\] (3.29)

\[
h_t(-1) = \begin{cases} P_Y(Y_t) - h_t(0), & Y_t < 0 \\ 0, & Y_t \geq 0 \end{cases}
\] (3.30)

\( P_Y(Y_t) \) and \( P(Y_t \mid c = 0) \) can be estimated from the data \( Y \) using non-parametric kernel estimation. \( P_Y(Y_t) \) is estimated using the full time segment, whereas \( P(Y_t \mid c = 0) \) considers only the pre-stimulus segment.

The actual classification procedure for significative synchrony changes is performed (for each frequency \( \omega \) and electrode pair \((e_1, e_2)\)) as follows:

1. Estimate the pre-stimulus distribution \( P_0(Y) \) and the full distribution \( P_Y(Y) \) using kernel density estimation with bandwidth given by Silverman’s rule of thumb [Silverman, 1986].

2. Estimate \( \alpha_0 = P(c = 0) = P_0(0)/P_Y(0). \)

3. For each \( t \), calculate \( h_t(k) \) for \( k = -1, 0, 1 \) as given by Equations (3.28, 3.29, and 3.30).

4. Normalize \( h_t \) to obtain \( \tilde{p}_t. \)
5. Obtain $p(k)$ by convolving $\hat{p}(k)$ with a Gaussian kernel $g$. The width $\sigma_g$ of the kernel controls the granularity of the $c$ field (see below).

6. Approximate the MPM estimator by $c_t = \text{argmax}_k p_t(k)$ for all $t$.

### 3.4.4 Granularity and choice of $\sigma_g$

When approximating $p(k)$, we need to use an adequate width $\sigma_g$ for the Gaussian kernel to filter out shorter homogeneous segments, which may not be significant in terms of persistence, and would instead break a longer segment. In particular, one would like that only those synchronous episodes with a duration greater than a minimum (physiologically motivated) value should be considered significant. According to studies related, for example, with the formation of short term memory [Jensen et al., 1996; Burle et al., 2000], this minimum value is in the order of one gamma oscillation, that is, around 20-50 ms.

To achieve this, one can define a granularity function $G(\sigma_g)$ for a given EEG dataset and see how it behaves with respect to $\sigma_g$. Our choice for $G$ is defined as follows: for a class time series $c_{\omega,e_1,e_2} = \{c_{\omega,e_1,e_2}(t)\}$ we define the number of homogeneous segments $n_{\omega,e_1,e_2}$ as

$$n_{\omega,e_1,e_2} = \sum_{t=T_s+1}^{T} 1 + [1 - \delta (c_{\omega,e_1,e_2}(t) = c_{\omega,e_1,e_2}(t-1))],$$

(3.31)

where $\delta(x)$ is the Kronecker delta function. Note that the sum is taken only on the post-stimulus segment. This is because we consider the pre-stimulus to be a single segment with class $c = 0$. 

---

Figure 3.3: Granularity functions $G_w(\sigma_g)$ for $w = 10, 15, 20, 25$ ms. We are interested in the lowest value of $\sigma_g$ for which $G_w(\sigma_g) < 0.05$. 

---

44
Table 3.1: Values of $\sigma_g$ (within a resolution of 0.5 ms) required to obtain $G_w(\sigma_g) < 0.05$, for various values of the granularity length $w$.

\[
\begin{array}{|c|c|}
\hline
w \text{ (ms)} & \sigma_g \text{ (ms)} \\
\hline
10 & 4 \\
15 & 7.5 \\
20 & 11.5 \\
25 & 16.5 \\
50 & 76 \\
\hline
\end{array}
\]

We also define $m_{\omega,e_1,e_2}(w)$ as the number of homogeneous segments whose length is less than $w$ (the desired granularity length). The granularity function is then given by

\[
G_w(\sigma_g) = \mathbb{E} \left[ \frac{m_{\omega,e_1,e_2}(w)}{m_{\omega,e_1,e_2}} \right],
\]

where the expected value is estimated across all frequencies and electrode pairs. $G_w(\sigma_g)$ estimates the probability of having segments with length shorter than $w$ for a given value of $\sigma_g$. One can then specify a value for $w$ and find $\sigma_g$ such that, for example, less than 5% of the segments have length less than $w$ (i.e., $G_w(\sigma_g) < 0.05$). Figure 3.3 shows the granularity function graphs for $w = 10, 15, 20, 25$ ms for the Figures experiment using the MPD measure.

Table 3.1 shows the value of $\sigma_g$ required for having less than 5% segments with length less than $w = 10, 15, 20, 25, 50$ ms. One can see that for values of $w$ greater than 25-30 ms, the required $\sigma_g$ grows considerably; however, the resulting label field $c$ does not seem to be seriously affected, even if relatively large values of $\sigma_g$ are used. In other words, the Bayesian method shown above seems to be very robust with respect to the kernel width $\sigma_g$. As an example, Table 3.2 shows the correlations, for the Figures experiment, between pairs of class fields estimated with different values of $\sigma_g$ (using the MPD measure). All correlations corresponding to values of $\sigma_g$ between 4 and 16.5 ms (which correspond to values of $w$ between 10 and 25 ms) are greater than 0.97, and correlations involving $\sigma_g = 76$ ms ($w = 50$ ms) are greater than 0.91.

These results suggest that one can loosely choose $\sigma_g$ from a relatively wide range of values (roughly between 10 and 50 ms). In particular, we have used $\sigma_g = 25$ ms for all our tests, and all the experiments.

### 3.4.5 Comparison between Bayesian estimation and thresholded p-values

We have estimated, for the MPD and CPPD measures and for each experiment, the threshold which maximizes the correlation between the class label field obtained from the Bayesian method and the one obtained from thresholded
Table 3.2: Correlations between pairs of class fields with different values of $\sigma_g$ (shown in bold typeface).

<table>
<thead>
<tr>
<th></th>
<th>4 ms</th>
<th>7.5 ms</th>
<th>11.5 ms</th>
<th>16.5 ms</th>
<th>76 ms</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 ms</td>
<td>1.000</td>
<td>0.991</td>
<td>0.983</td>
<td>0.974</td>
<td>0.912</td>
</tr>
<tr>
<td>7.5 ms</td>
<td>0.991</td>
<td>1.000</td>
<td>0.991</td>
<td>0.982</td>
<td>0.916</td>
</tr>
<tr>
<td>11.5 ms</td>
<td>0.983</td>
<td>0.991</td>
<td>1.000</td>
<td>0.990</td>
<td>0.921</td>
</tr>
<tr>
<td>16.5 ms</td>
<td>0.974</td>
<td>0.982</td>
<td>0.990</td>
<td>1.000</td>
<td>0.928</td>
</tr>
<tr>
<td>76 ms</td>
<td>0.912</td>
<td>0.916</td>
<td>0.921</td>
<td>0.928</td>
<td>1.000</td>
</tr>
</tbody>
</table>

Table 3.3: Threshold required to maximize the correlation between the thresholded p-values classification, and the Bayesian classification (with $\sigma_g = 16.5$ ms). The correlation is shown for each test experiment, and for the MPD and CPPD measures.

<table>
<thead>
<tr>
<th></th>
<th>Figures</th>
<th>Words</th>
<th>Go</th>
<th>NoGo</th>
<th>Stern-3</th>
<th>Stern-5</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPD Threshold</td>
<td>0.11</td>
<td>0.09</td>
<td>0.04</td>
<td>0.03</td>
<td>0.04</td>
<td>0.04</td>
</tr>
<tr>
<td>CPPD Threshold</td>
<td>0.06</td>
<td>0.06</td>
<td>0.01</td>
<td>0.01</td>
<td>0.02</td>
<td>0.02</td>
</tr>
</tbody>
</table>

This means that the Bayesian approach yields results roughly equivalent to thresholded p-values with a threshold between 0.01 and 0.1, but this value has to be adjusted for each experiment and it is not clear how this should be done, whereas the Bayesian method permits one to specify the granularity independently of the particular experiment.

The correlations shown above change very little with respect to the granularity value, which is expected, since the Bayesian method is barely sensitive to the granularity parameter. For example, for the Figures experiment with the MPD measure, granularity values of 10, 15, 20, and 25 ms yield roughly equivalent results to a p-value threshold of 0.11, while a granularity of about 50ms corresponds to a threshold of 0.10. This robustness with respect to the granularity parameter is one of the reasons we chose the Bayesian approach.

### 3.5 Visualization

Synchrony data is in a four-dimensional space. Because of this, the visualization problem is not a trivial one, especially since one would like to keep as much detail as possible while displaying the data. It is very common to either focus on specific points in the time-frequency plane or condense the data by averaging in one or more dimensions.
Figure 3.4: Multitoposcopic displays for a 120-channel dataset using the MPD measure: (a) uses only channels Fp1, Fp2, F3, F4, C3, C4, P3, P4, O1, O2, F7, F8, T3, T4, T5, T6, Fz, Cz, and Pz from the 10/20 system: red regions represent electrode pairs that show a significant increase in synchrony ($c = 1$), while green regions correspond to significant synchrony decreases ($c = -1$). Graphs (b) and (c) use the 120 channels grouped in 19 cortical areas corresponding to the Voronoi partition whose centers are the electrodes used in (a). For each area, one can plot a toposcope representing a statistic of the synchrony classes between each of the 120 electrodes and the electrodes within the area: (b) uses the average class $\bar{c}$ (which may be between -1 and 1), whereas (c) uses the class mode $\hat{c}$ (see text for details). The 3 multitoposcopes represent the SP obtained at $t=515$ ms and $f=11$ Hz during a Go/NoGo experiment (see [Harmony et al., 2006] for details); the rest of the examples presented in this paper correspond to the Figures experiment described in the text.

For a fixed time $t$ and frequency $\omega$ it is possible to show the distribution of the synchrony pattern (SP) given by the class values $c_{\omega,e_1,e_2,t}$ in a multitoposcopic display, in which for each electrode $e_1$ one displays a head diagram (also called “toposcope”) - within a bigger head - that shows the distribution of $c$ across all sites $e_2$ (an example of this display can be seen in Figure 3.4a).

A problem arises when one deals with high electrode density recordings since only so many toposcopes can be displayed within the bigger head. One can use interpolation techniques to display high-density data in a single toposcope (in particular, we use a Voronoi partition [Aurenhammer, 1991] obtained by assigning each pixel in the toposcope the class corresponding to the nearest electrode), but the number of toposcopes may have to be reduced. One possibility consists in grouping the electrodes in $N_g$ cortical areas $\{G_1, \ldots, G_{N_g}\}$, using again a Voronoi partition whose centers are, for example, the sites Fp1, Fp2, F3, F4, C3, C4, P3, P4, O1, O2, F7, F8, T3, T4, T5, T6, Fz, Cz, and Pz of the 10/20 system. One can then plot a toposcope for each group $G_k$ computing, for each high-resolution electrode, the “most representative” synchrony class between this electrode and the electrodes in $G_k$. This most representative class may be computed in several ways; for example, the average class $\bar{c}$ of electrode $e$ to
group $G_k$, which is computed as:

$$
\bar{c}_{\omega,k,e,t} = \frac{1}{|G_k|} \sum_{e' \in G_k} c_{\omega,e,e',t}
$$

(3.33)

or the class mode $\hat{c}$, which is computed as:

$$
\hat{c}_{\omega,k,e,t} = \arg \max_{q \in \{-1,0,1\}} \left\{ \sum_{e' \in G_k} \delta(c_{\omega,e,e',t} - q) \right\},
$$

(3.34)

where $\delta$ is the Kronecker delta function. An example of multitoposcopes corresponding to $\bar{c}$ and $\hat{c}$ are shown in Figures 3.4b and 3.4c. More sophisticated methods could involve electrode clustering or TFT regularization; however, they are beyond the scope of this work.

Multitoposcopic displays are useful to show a detailed connectivity pattern for a fixed time and frequency; however, it is important to visualize larger regions of the time-frequency plane in order to localize zones of interest where the synchrony pattern remains almost constant and might be related to specific cognitive processes. Following the procedure given by [Marroquin et al., 2004] to analyze relative amplitude changes, we can use a Time-Frequency-Topography (TFT) display to present the data by reducing only one spatial dimension. We do this not by averaging but counting, for each electrode $e$, the number of electrodes that have significantly increased or decreased their synchrony with $e$.

In other words, we can build a \textit{synchrony increase histogram} (SIH) given by

$$
H^+_{\omega,e}(t) = \sum_{e' = 1}^{N_e} \delta(c_{\omega,e,e',t} - 1),
$$

(3.35)

where $\delta(x)$ is the Kronecker delta function. $H^+_{\omega,e}(t)$ is the number of significantly stronger couplings (with respect to the pre-stimulus segment) for electrode $e$ at time $t$ and frequency $\omega$. This can be thought as a degree of connectivity involving electrode $e$ relative to the pre-stimulus.

Similarly, we can define a \textit{synchrony decrease histogram} (SDH) as

$$
H^-_{\omega,e}(t) = \sum_{e' = 1}^{N_e} \delta(c_{\omega,e,e',t} - (-1)),
$$

(3.36)

These histograms can be presented in a TFT display as shown in Figure 3.5.

3.5.1 Partitioned TF synchrony maps

The TFT visualization system provides at the same time a detailed view that shows which regions of the cortex are involved in synchronous processes at
Figure 3.5: Synchrony histograms for the Figures experiment displayed as Time-Frequency-Topography (TFT) maps. The left and right maps show the synchrony increase and decrease histograms, respectively, for the MPD measure. The color scale represents the number of electrodes whose synchronization with a given electrode $e$ increases (left) or decreases (right) significantly.

It is also possible to produce an automated partition by frequency bands (delta, theta, alpha, etc.), and at regular time intervals. An example of this is shown in Figure 3.7. The frequency bands are delta (1 to 3 Hz), theta (4 to 7 Hz), alpha (8 to 12 Hz) and low beta (13 to 18 Hz) with a segment interval of 300 ms. In the lower-left corner of each segment we show the estimated variance of $c_{\omega,e,e',t}$. For regions with high variance, the representative SP may not be consistent across region; however, one can subdivide the region for a more detailed analysis.
Figure 3.6: Manually-segmented MPD synchrony map for the Figures experiment. The left map uses the average class (represented by the color scale) for each electrode pair to construct the representative SP’s, whereas the right map uses the class mode.

Figure 3.7: Automatically partitioned MPD maps for the Figures experiment (using the average class of each TF region as representative). Regions with high-variance can be subdivided into smaller, more consistent regions.
3.6 Gabor filters vs Sinusoidal filters

We have performed a series of tests in order to demonstrate the improvement of SQF’s over Gabor filters for synchrony estimation. From Figure 2.1 one can see that the distortion produced by the Gabor filter (at tuning frequencies less than 6 Hz) consists of an oscillation at twice the tuning frequency. For these tests we applied the synchrony measure (based on $\Delta \Phi$) and statistical analysis described below. Using Gabor filters, the sensitivity of the synchrony measure shows a cyclic dependence on the time $t$, which corresponds to the distortions in the phase observed in Figure 2.1.

The simulated data used for the tests is obtained from a very simple model which consists of a 2 Hz sinusoidal plus noise. The signals are given by:

$$V_{j,e} = \cos(2\pi f_0 t + \phi_{j,e}) + R_{j,e}(r,t), \quad \phi_{j,e} \sim \mathcal{N}(0, \sigma),$$

(3.37)

where $j$ is the trial number, $e$ the electrode index, $f_0 = 2$ Hz is the frequency of the sinusoidal, and $\sigma$ is the standard deviation of the distribution of the phases, which controls the degree of synchrony between the signals. The noise function $R_{j,e}(r,t)$ depends on the “noise level” $r$. For our tests we used two electrodes ($e = 1, 2$), $\sigma_{\text{pre}} = \pi/10$ (pre-stimulus $\sigma$), and $\sigma_{\text{post}} = \pi/20$ (post-stimulus $\sigma$). Since $\sigma_{\text{pre}} > \sigma_{\text{post}}$, we are thus modeling an increase of synchrony in the full post-stimulus segment. We have generated 20 datasets with 50 trials each. For each dataset we perform the procedure described below in order to obtain a label field $c_{\omega, e_1, e_2}(t)$, which indicates if synchrony between $e_1$ and $e_2$ at a frequency $\omega$ is significantly higher ($c = 1$), lower ($c = -1$), or equal ($c = 0$) with respect to the pre-stimulus average. Focusing only on frequency $\omega = 2\pi f_0$ and electrode pair $< 1, 2 >$, we expect to obtain $c(t) = 1$ for all $t$ in the post-stimulus.

For the first test we used Gaussian noise given by

$$R_{j,e}(r,t) \sim \mathcal{N}(0, r).$$

(3.38)

With a noise level $r = 0.5$, the average class $\bar{c}(t)$ across all datasets is shown in Figure 3.8 (a). The blue line represents the results obtained using Gabor filters, whereas the red line corresponds to sinusoidal quadrature filters. The oscillating distortion is clearly present with the Gabor filters, and it corresponds to a frequency of 4 Hz (twice the tuning frequency). Figure 3.8 (b) shows the proportion of detected couplings ($c(t) = 1$) in the post-stimulus segment with respect to the noise level $r$ - these results are also averaged across the 20 datasets. The proportion should remain close to 1, however, Gabor filters clearly start to fail with $r > 0.3$.

The second test uses a sum of interfering sine waves as noise function:

$$R_{j,e}(r,t) = \sum_{f \neq f_0} a_f \cos(2\pi f_t + \phi_f), \quad a_f \sim \mathcal{U}(0, r), \quad \phi_f \sim \mathcal{U}(0, 2\pi).$$

(3.39)

The sum is taken across frequencies from 1 to 50 Hz, except the frequency of interest $f_0$, and the components are attenuated according to their frequency.
Figure 3.8: Comparison between sinusoidal and Gabor filters using simulated data with Gaussian noise for (a) and (b), and with interference of sinusoidals for (c) and (d). Graphs (a) and (c) show the average class $\bar{c}(t)$ computed across 20 datasets, whereas (b) and (d) show the proportion of detected couplings with respect to the noise level.

As in the previous test, we calculated the average class $\bar{c}$ and the proportion of detected couplings with respect to the noise level. The results, presented in Figure 3.8 (c) and (d), show the same differences as in the previous example.

It is worth mentioning that using real EEG datasets (such as the Figure experiment signals), there are clear differences between the results obtained with SQF’s and those obtained from Gabor filters in the first 6 Hz. The tests with simulated data indicate that the results with Gabor filters may not be reliable in the delta and theta bands.

### 3.7 Reducing volume conduction effects

In Section 1.6 we discussed the problem of volume conduction in EEG, which may introduce spurious correlations in the EEG signals. One of the best solutions so far, as reported by Nunez et al. [1995; 1997; 2000], consists in estimating the Surface Laplacian (SL) of the raw potentials; however, a high spatial sampling rate (e.g., more than 64 electrodes distributed across the whole surface) is required to obtain an accurate SL, and even in that case, the phase of the SL signals may be very sensitive to noise [Junghöfer et al., 1999; Hagemann et al.,]
2001]. On the other hand, most of our datasets contain only 19 or 20 channels, which is not enough for a reliable SL estimation.

Another solution, developed in collaboration with Joaquín Peña, is to use instead the phase of a new set of signals which are projections of the scalp potentials to cortical sites located directly below the actual electrodes. These signals may be obtained by placing dipoles (sources) in the cortical sites and solving the corresponding inverse problem.

The relation between the potentials and the current density at each source is given by:

$$\Phi = KJ,$$

where $\Phi$ is a vector (of length $N_e$) of potentials measured at each electrode, $J = (j_1, \ldots, j_{N_s})$ is comprised of the current densities $j_k = (j_{kx}, j_{ky}, j_{kz})^T$ at each source $k = 1, \ldots, N_s$, and $K$ is an $N_e \times 3N_s$ matrix (the lead field), which transforms current densities into surface potentials [Malmivuo and Plonsey, 1995; Pascual-Marqui, 1999]. $K$ depends on the electrode positions, the physical and geometrical properties of the volume conductor, and the position of the source dipoles. In particular, we compute $K$ using a three concentric sphere head model [Zhang, 1995].

The inverse problem consists in finding $J$, given $\Phi$ and $K$. This is typically an ill-posed problem, since the number of unknowns ($3N_s$) is usually larger than the number of equations ($N_e$). In order to make this problem well posed, we place one dipole in the cortex area directly below each electrode, with radial direction (i.e., perpendicular to the surface and pointing outwards).

In general, one can write the current densities for each source as $j_k = \alpha_k q_k$, $|q_k| = 1$, where $q_k$ is the direction of the $k$-th dipole, and $\alpha_k$ its strength. One can also write the lead field $K$ as:

$$K = \begin{bmatrix} k_{11} & k_{12} & \cdots & k_{1N_s} \\ k_{21} & k_{22} & \cdots & k_{2N_s} \\ \vdots & \vdots & \ddots & \vdots \\ k_{N_e1} & k_{N_e2} & \cdots & k_{N_eN_s} \end{bmatrix},$$

with $k_{ij} = [k_{i(3j)} \ k_{i(3j+1)} \ k_{i(3j+2)}]$.

The surface potentials are thus given by

$$\Phi = \begin{bmatrix} k_{11}\alpha_1 q_1 + k_{12}\alpha_2 q_2 + \cdots + k_{1N_s}\alpha_{N_s} q_{N_s} \\ k_{21}\alpha_1 q_1 + k_{22}\alpha_2 q_2 + \cdots + k_{2N_s}\alpha_{N_s} q_{N_s} \\ \vdots \\ k_{N_e1}\alpha_1 q_1 + k_{N_e2}\alpha_2 q_2 + \cdots + k_{N_eN_s}\alpha_{N_s} q_{N_s} \end{bmatrix}$$

(3.42)
Figure 3.9: (a) Dipole groups under each electrode within a three concentric sphere model. (b) Scalp potentials obtained with the forward equation and the given group strengths.

If the dipole directions $q_k$, $k = 1, \ldots, N_s$ are known, then one can compute the matrix $L$. Moreover, if there is only one source per electrode (i.e., $N_s = N_e$), then the problem becomes well-posed as the number of unknowns (i.e., the $\alpha_k$'s) equals the number of equations. In this case, $L$ is square and the inverse problem has a unique solution given by:

$$\hat{\alpha} = L^\dagger \Phi.$$  

(3.43)

Where $L^\dagger$ is, in general, the pseudoinverse of $L$ [Vogel, 2002], which can be obtained, for example, by singular value decomposition (SVD - [Press et al., 1992]).

Instead of a single dipole per electrode, one can also place a group of cortical dipoles (for each recording site) whose strengths decrease exponentially with the distance to the central dipole (the one located right below the electrode - see Figure 3.9a). In other words, the amplitude of all the dipoles in the group is determined by the strength of the central dipole (the “group strength”). The inverse problem can be solved in a similar manner: one can write the potentials $\Phi$ as the product of a square matrix $L$ and the vector of group strengths $\alpha$, where $L$ (and its pseudoinverse) can be computed. Figure 3.9b shows an example of
scalp potentials obtained from the given group strengths, using the forward model \( \Phi = L\alpha \).

The projection of scalp potentials to the cortical group strengths reduces the conductor effects of the scalp and skull, and provides better spatial localization than the raw scalp potentials, but with a magnitude large enough to obtain reliable phase estimations. Figures 3.10a and 3.10b show the resulting synchrony patterns obtained for synthetic signals using raw potentials (3.10a) and the projected signals (3.10b). The test signals were obtained by placing one dipole near F3 and another between P4 and T6 of the 10/20 system, whose output is a sinusoidal signal, and solving the forward problem to obtain scalp readings. Noise was added to all electrode readings. The dipoles are unsynchronized (out of phase) in the pre-stimulus segment (not shown) and in perfect synchrony in the post-stimulus. The graphs show significant increment (red) or decrement (green) in synchrony for all electrode pairs. The synchrony pattern obtained from projected signals clearly shows a better localization of the synchronized regions. Another example, with real EEG signals from the Figures experiment, is shown in Figures 3.10c (scalp potentials) and 3.10d (cortical projections). Note that, in this case, synchronous activity also appears to be smeared when using potentials: sites P4 and Oz show a significant increment in synchrony with almost every other site; however, most of these couplings do not show when using projected signals, possibly indicating that they were spurious.
Chapter 4

Comparative study between synchrony measures

4.1 Measure comparisons in the literature

A few works have focused on the evaluation of different synchronization measures, using real and simulated EEG data. In one of these papers ([Quian-Quiroga et al., 2002]), interaction between the left and right hemispheres of a rat are analyzed with various synchrony measures. This includes both linear (correlation / coherence) and non-linear (generalized synchronization, mutual information, phase-locking) interdependences.

According to Quian-Quiroga, all the measures (with the exception of MI) ranked the synchronization levels in the same way; that is, the periods of low and high synchrony, respectively, were similarly distinguished by every measure. However, the separation between low and high synchrony periods was more noticeable with non-linear measures (GS and STPLS) than with coherence. This may indicate that most couplings have a non-linear nature (e.g. phase-locking, as Varela suggests). In the case of mutual information, Quian-Quiroga et al. used a time-embedding technique to transform the data to a $m$-dimensional space, which made it difficult to obtain a good estimation of the joint entropy as the joint probability matrix was very sparse. Because of this, it was not possible to obtain robust estimations of the MI, which dramatically depended on $m$.

In 2003, David and Friston introduced a neural mass EEG model, which simulates the activity of populations of neurons, under the assumption that the state of the population can be approximated with very few state variables [David and Friston, 2003b]. This model is based on a simpler model ([Jansen and Rit, 1995]) that is able to produce any of the common EEG rhythms (alpha, beta, etc.).

In the simulations performed by David et al., it was observed that when both
areas were bidirectionally coupled, the phase-shift between the two signals was equal to zero, and it was shown that for bidirectional couplings, the phase-shift is equal to either 0 or \( \pi \), depending on the propagation delay. According to David and Friston [2003b] and Varela et al. [2001], the proportion of reciprocal connections in the brain is very high. This supports the findings in [Friston et al., 1997; Rodriguez et al., 1999], where a tendency of the phase-differences towards zero were associated with synchronous episodes (see Sections 2.3.1 and 2.3.2).

In their following article ([David et al., 2004]), David and Friston tested various synchronization measures using simulated data. The measures were: cross-correlation (with zero time-lag), mutual information, generalized synchronization, and the single-trial phase-locking value. The simulations were obtained with their neural mass model, using a symmetric configuration for a bidirectional coupling, and a parameter \( k \in [0, 1] \), which specifies the coupling strength. A series of surrogate (uncoupled) signals were generated with \( k = 0 \) in order to obtain the null distributions of the synchrony measures.

In a first analysis, the coupling sensitivity of the various measures was tested using broadband signals, as a function of the coupling strength \( k \). While all measures increased accordingly with \( k \), GS showed a better response to weak couplings (\( k < 0.125 \)), while MI was better for strong couplings. The next analysis evaluates the response to the time-window size used to estimate the measures. The window was varied from 0.2 to 2 s, and the z-scores (with respect to the null distribution from an uncoupled model) were estimated for each measure. In this case, GS resulted to be the most reliable estimator.

For narrow-band analysis, the coupled and uncoupled signals were band-pass filtered from 2 to 50 Hz, each 2 Hz, and the synchrony measures were estimated for each frequency. The resulting z-scores were plotted as a function of frequency. The graphs for all measures had a similar shape; however, the z-scores for mutual information were much higher, suggesting that MI may be more sensitive to frequency-specific synchrony.

The next analysis focused on the sensitivity of the measures to non-linear couplings. In this case, the surrogate population is designed to preserve the linear properties of the original data, while destroying the non-linear dependencies (see [David et al., 2004] for details). This method was applied to the simulated broadband signals, and the synchrony measures were estimated as a function of the coupling strength. Cross-correlation and mutual information were insensitive to non-linear couplings, while with GS and phase synchronization, the non-linearities accounted for a small fraction of the synchrony value.

David et al. concluded that although the results of the four measures tested were very similar, one should be careful when studying functional connectivity using electrophysiological signals. One should not make too many assumptions about the type of interactions; therefore, instead of relying on a single measure, it is more advisable to apply various measures, each of which is sensitive to different aspects of synchronization.
4.2 Comparison of synchrony measures

The procedure described in Chapter 3 can be used with any of the synchrony measures presented in Section 2.3. In particular, we have tested the PLS, STPLS, and coherence measures. GS and MI have not been implemented for the reasons explained in Sections 2.3.4 and 2.3.5.

For the rest of this chapter, we will present various interesting results which may help in determining the advantages and disadvantages of each measure.

4.2.1 TFT Synchrony maps

We have computed the following measures for the Figures experiment:

- Mean Phase Difference (MPD)
- Cumulative Probability of Phase Difference (CPPD), with $\epsilon = \pi/5$ rad.
- Lachaux’s Phase Locking Statistic (PLS)
- Single-Trial PLS (STPLS), with a window size of 105 ms ($w = 10$ samples).
- Coherence, with a window size of 105 ms.

Figure 4.1 show the SIH’s for all the measures (full SIH, SDH, and partitioned maps will be presented in Appendix ??).

From these maps, one can observe that CPPD and PLS produce similar results to the MPD measure. On the other hand, STPLS and coherence yield similar results themselves but different from MPD. This suggests that the synchrony measures may be divided in two groups.

4.2.2 Correlation between measures

Given two synchrony measures, $\mu^1$ and $\mu^2$, one can estimate the class label fields $c^1$ and $c^2$, and compute a correlation measure between the two class fields. Typically, the correlation between two random variables $X$ and $Y$ is given by

$$\text{cor}(X, Y) = \frac{E[(X - EX)(Y - EY)]}{\sigma_X \sigma_Y},$$

(4.1)

where $E[\cdot]$ denotes the expected value, and $\sigma_X$ is the standard deviation of $X$. Using this correlation measure, we have computed the degree of interdependence between each pair of measures for the Figures experiment. The results are presented in Table 4.1, where two groups are clearly defined by high correlation values: one group consisting of MPD, CPPD, and PLS, and the other consisting of STPLS and coherence. Similar results have been obtained with
Figure 4.1: Synchrony increase histograms for the Figures experiment using the five synchrony measures: MPD, CPPD, PLS, STPLS, and coherence. The bottom-right graph corresponds to the TFT map of significant changes in the LPC measure (red represents significant increases, whereas green corresponds to significant decreases) - see text for details.
Another dependence measure, which may be more adequate for class data, can be defined as the probability of equal class - PEC, which can be estimated as
\[ \text{pec}(\mu_1, \mu_2) = \frac{1}{N_f N_e N_t} \sum_{\omega, e_1, e_2, t} \delta(c^1_{\omega, e_1, e_2, t} - c^2_{\omega, e_1, e_2, t}) \],
(4.2)
where \( \delta \) is the Kronecker delta function. Table 4.2 shows the dependence values for all pairs of measures using Equation 4.2; the two groups of measures can also be distinguished here.

It is worth noting that STPLS and coherence are computed across a time window, whereas the other measures are “instantaneous”, in the sense that they only use the phase difference at a single time point. This may be a determining factor in the behavior of a given measure.

One interesting fact is the high correlation between MPD and PLS. By looking at Equation (2.15) one can see that PLS actually measures the consistency of the phase difference across all trials. A high correlation between MPD and PLS suggests that the processes that result in high in-phase synchronization are fairly consistent across trials and subjects, and thus may be related to the task.

Furthermore, the CPPD measure is also highly correlated to the MPD, suggesting that synchronization effectively happens with near-zero phase difference.
This is in accordance with the neural mass model proposed by David and Friston [2003a] and other works ([Friston et al., 1997; Rodriguez et al., 1999]) which have also found zero-centered phase difference distributions during synchronous episodes between two electrodes.

4.2.3 Response to induced activity

One could argue that instantaneous synchrony measures (such as MPD) may be more sensitive to variations in the latency of event-related responses across trials (latency jitter); in other words, these measures may not respond adequately to induced activity.

Typically, induced activity is revealed by performing a time-frequency decomposition of each trial, and averaging a positive definite function of the filters’ output (e.g., the power) across all trials, whereas evoked activity is obtained by first averaging the raw signals across trials, and then applying a positive definite function (power analysis) to the average signals [David et al., 2006]. The difference is that in the former case (induced responses) there will be no cancelations during averaging, whereas in the latter (evoked) case, there may be cancelations due to differences in sign across trials. It is clear then, that the methodology presented in Chapter 3 is basically an induced response analysis, where the positive definite function is the synchrony measure (we may call it “induced synchrony”), instead of the power; this function is averaged across trials and then the baseline is subtracted. This suggests that the analysis procedure, with any of the synchrony measures previously presented, should be at least as sensitive to induced responses as the typical average-power analysis. Of course, because of additional averaging across time, STPLS and coherence may be more robust to larger amounts of latency jitter.

In Section 6.4.1 we will briefly discuss some preliminary results about the relationship between evoked potentials and induced synchrony, and the mechanisms which generate both types of activity.

4.2.4 Local phase dispersion

Let us recall the equation for Lachaux’s single-trial phase-locking statistic:

$$\text{STPLS}_{e_1,e_2}(t) = \frac{1}{2w + 1} \left| \sum_{t'=t-w}^{t+w} \exp[i(\phi_{e_1}(t') - \phi_{e_2}(t'))] \right|.$$  (4.3)

The STPLS is equal to one minus the dispersion of the phase difference between the two electrodes under study, estimated across a time window centered at $t$. Note that if one of the phases remains relatively constant within the time window, then the STPLS will be approximately equal to one minus the dispersion of the other phase. This suggests that it may be interesting to study the correlation between the local phase dispersion and each synchrony measure. To
Table 4.3: Correlation between significance indexes of LPC/Power changes, and the average class at each electrode, for each synchrony measure.

<table>
<thead>
<tr>
<th></th>
<th>MPD</th>
<th>CPPD</th>
<th>PLS</th>
<th>STPLS</th>
<th>Coherence</th>
</tr>
</thead>
<tbody>
<tr>
<td>LPC</td>
<td>0.091</td>
<td>0.035</td>
<td>0.131</td>
<td>0.698</td>
<td>0.673</td>
</tr>
<tr>
<td>Power</td>
<td>-0.028</td>
<td>-0.122</td>
<td>0.067</td>
<td>0.266</td>
<td>0.273</td>
</tr>
</tbody>
</table>

do this, one can define a local phase constancy (LPC) measure as the average across trials of one minus the dispersion of the phase at each electrode. The LPC is thus given by

$$\text{LPC}_{\omega,e}(t) = \frac{1}{N_r} \sum_{j=1}^{N_r} \frac{1}{2w+1} \sum_{t' = t-w}^{t+w} \exp(i\phi_{j,\omega,e})$$

(4.4)

One can then estimate a TFT map of significant changes of the LPC measure with respect to the pre-stimulus, as described in Section 3.4. In particular, we use the significance indexes obtained from p-values (Section 3.4.1), which we denote here as LPC$^S_{\omega,e}(t)$. The TFT map of LPC changes is shown in Figure 4.1 (bottom-right). Note the similarity between this map and the SIH’s for STPLS and coherence: increases in the LPC correspond to increases in STPLS and coherence; moreover, the green region in the LPC map (around 9-12 Hz) corresponds to high values in the SDH’s of STPLS and coherence (see Appendix ??).

In order to estimate the correlation between significant changes in the LPC and significant changes in a synchrony measure, one can compute the average class at each electrode, which is given by

$$c_{\omega,e}^{\text{avg}}(t) = \frac{1}{N_e - 1} \sum_{e' = 1}^{N_e} c_{\omega,e,e',t}.$$  

(4.5)

Here we divide by $N_e - 1$ instead of $N_e$ since $c_{\omega,e,e,t} = 0$ for every electrode $e$. Table 4.3 shows the correlation between LPC$^S$ and the average class ($c_{\text{avg}}$) corresponding to each of the tested measures. Note that the correlation values corresponding to STPLS and coherence are significantly higher than for the other measures.

These results suggest that STPLS and coherence may be seriously influenced by changes in the dispersion of the local phase measured at each electrode; in other words, they may be more sensitive to local processes which are not necessarily related to the synchronization between the two electrodes under study. For these reasons, we believe that instantaneous measures may be better suited for synchrony analysis.

As a sidenote, Table 4.3 also shows that STPLS and coherence are slightly more correlated to instantaneous power changes. This makes sense for coherence, since it uses both the magnitude and phase of the signals; however, the
correlation indexes are not large enough to suggest a strong dependence between power and long-range synchrony.

4.2.5 Influence of volume conduction

We have already discussed the importance of the volume conductor in the study of EEG synchrony. Some authors argue that a synchronization measure that favors phase differences of zero (such as the MPD) would be highly sensitive to volume conduction; however, this is not entirely clear. Lachaux et al. [1999] mention that “another common assumption is that the phase difference between electrodes should be zero in case of conduction synchrony. This is usually false: even if two electrodes record the same group of sources, the signals of these electrodes are different linear combinations of the source amplitudes, ... Therefore, the phase of the original signals recorded by the two electrodes should be different, except if all the sources have the same phase”. In other words, if several sources influence two different sites via volume conduction, their net influence on the resulting electrode signals will correspond to different linear combinations of these source signals (with coefficients given by the corresponding attenuations), thus producing different phases. From this viewpoint, there is no reason to think that MPD or CPPD will be more influenced by volume conduction than the other measures. Furthermore, the MPD and CPPD measures are highly correlated to Lachaux’s PLS measure, which measures synchrony with a constant phase lag (not necessarily zero), suggesting that at least these three measures are similarly affected by volume conduction.

Lachaux also suggests that one may be able to identify conduction synchrony by looking at the neighboring sites of the two electrodes. In other words, if conduction synchrony is observed between two electrodes, then one must also observe high synchrony between their neighbors. One can test this by defining a neighborhood $N_i$ for each electrode $e_i$, and then counting, for each electrode pair $<e_1, e_2>$ for which an increase in synchrony is observed, the number of pairs $<e_r, e_s>$, $e_r \in N_1$, $e_s \in N_2$ whose synchrony also increases. If this number is relatively high for all $t$, it may suggest a higher amount of spurious couplings due to volume conduction (although this is not necessarily true). One can then compare these values for different synchronization measures to see if any of them may be more sensitive to conduction effects. With this in mind, one can define a neighbor-synchrony measure $Z^+$, which is given by:

$$Z^+_{\omega, e_1, e_2, t} = \frac{\sum_{r \in N_1, s \in N_2} \delta(c_{\omega, e_r, e_s, t} - 1) \cdot \delta(c_{\omega, e_1, e_2, t} - 1)}{|N_1||N_2|},$$

where $c$ is the class label field obtained from thresholded p-values (as shown in Section 3.4.1). Here we avoid using the Bayesian classification since the regularization may introduce spurious correlations among neighboring electrodes. Note also that we are dividing by the size of the neighborhoods; this is done in order to normalize the measure, since different electrodes may have differ-
Figure 4.2: TFT maps of neighbor synchrony ($Z^+$) for the Figures experiment using the five synchrony measures: MPD, CPPD, PLS, STPLS, and coherence.

ent neighborhood sizes. We have chosen a neighborhood radius such that each electrode has at least one neighbor.

By inspecting the TFT maps of the average $Z^+$ for each electrode (Figure 4.2), one can see that, if anything, STPLS and coherence show larger $Z^+$ values, although in a narrower frequency band, than the other measures. In order to obtain a more conclusive result, we averaged $Z^+$ across all the electrode pairs which show an increase of synchrony, and across all time and frequency points. The averaged $\overline{Z^+}$ is thus given by

$$\overline{Z^+} = \frac{\sum_{\omega,e_1,e_2,t} Z_{\omega,e_1,e_2,t} \delta(c_{\omega,e_1,e_2,t} - 1)}{\sum_{\omega,e_1,e_2,t} \delta(c_{\omega,e_1,e_2,t} - 1)}.$$ (4.7)

One can think of $\overline{Z^+}$ as the conditional probability of some neighbor of $e_1$ increasing its synchrony with some neighbor of $e_2$, given that the synchrony between $e_1$ and $e_2$ increases. Table 4.4 presents the resulting $\overline{Z^+}$ for each synchrony measure. Effectively (and surprisingly), STPLS and Coherence show a higher average neighbor synchrony, although the difference does not seem significant. In any case, it is clear that the in-phase synchrony measures (MPD and CPPD) do not yield $\overline{Z^+}$ values which are significantly greater than the other measures, and thus there is no evidence supporting the claim that these measures may be more sensitive to volume conduction effects.
Table 4.4: Average neighbor synchrony $Z^+$ for each synchrony measure applied to the Figures experiment.

### 4.2.6 Conclusions

The results shown in the previous sections suggest that

1. Phase synchrony effectively happens, in most cases, with near-zero phase difference between pairs of leads.
2. Phase synchrony, when happens, is relatively consistent across trials.
3. Instantaneous phase-lock measures are less sensitive to changes in the dispersion of the local phase at each electrode, and thus may be better suited for synchrony analysis than measures which are computed across a time window (disregarding filtering effects).
4. All measures should respond to induced activity; however, measures computed across a time window may be less sensitive to latency jitter.
5. All measures seem to be equally sensitive to volume conductor effects.
Chapter 5

Apparent phase and amplitude

In the previous chapters we have described our methodology for the analysis of synchronization patterns, and shown the advantages of the proposed measures (MPD and CPPD) over some of the measures in the literature. We have used these techniques to study a few real EEG datasets, and now we proceed to describe our findings.

In this chapter, we propose a simple mathematical model of the narrow-band signal captured by an electrode. With this model, one can explain and simulate most of the synchronization patterns that result from the analysis of the experiments. The following chapter will deal with the neurophysiological interpretation of such patterns.

5.1 Motivation

From the analysis of the Figures, Words, Go/No-Go, and Sternberg experiments, one can observe certain phenomena (some of which have been previously reported) that may appear counter-intuitive; for example:

1. Synchronous episodes occur with near-zero phase lag, regardless of the distance between leads. This has also been observed in [Friston et al., 1997] and [Rodriguez et al., 1999], and is in accordance with David and Friston’s neural mass model.

2. It is possible that two or more electrodes increase their synchrony with another electrode, which acts as a nodal point, and at the same time observe a decrease of synchronization between the non-nodal sites. One example of this phenomenon is the synchrony pattern observed in the alpha band (8-12 Hz) in Figures 3.6 and 3.7: the occipital leads increase their synchrony with most of the other electrodes, while fronto-parietal
leads show a decrease of synchrony among them. This phenomenon can also been observed in other studies, such as [Rodriguez et al., 1999], [Varela et al., 2001], and [David et al., 2003b].

3. When an electrode $e$ increases its synchrony with many other sites, there is a high probability of observing a significant power change at $e$ in the same frequency band. If the non-nodal sites show a decrease of synchronization, the power at $e$ will decrease, and vice versa.

These phenomena may be explained if one supposes that the signal recorded by a scalp electrode results in fact from the sum of macroscopic oscillations produced by functionally different neural populations. This assumption is supported by the following considerations: a scalp electrode provides estimates of synaptic action averaged over tissue masses containing between $10^7$ and $10^9$ neurons [Nunez, 1995]. There is evidence (obtained, for example, using retrogradely transported fluorescent dyes [Morecraft et al., 1993], and single-cell recordings [Quintana and Fuster, 1999]) of extensively intermingled populations of neurons connected with different areas (e.g., frontal eye fields and posterior parietal cortex). Moreover, when performing a computational task, neurons may quickly associate into a functional group while disassociating from concurrently activated groups [Haalman and Vaadia, 1998].

5.2 Population model

The facts exposed above suggest that one may be able to model certain aspects of the EEG dynamics by expressing each electrode signal as the sum of oscillating functions (e.g., sinusoidals) corresponding to the average activity of each contributing population. It is also plausible that, in order to develop a macroscopic oscillation, the neurons of each sub-population must be synchronized. Therefore, a reasonable model for a complex narrow-band EEG signal is:

$$V(t) = \sum_k \alpha_k \cos(\omega_k t + \phi_k), \quad (5.1)$$

where the positive coefficients $\alpha_k$ depend on the amplitude and relative size of subpopulation $k$, and $\phi_k$ is the corresponding phase. Note that this model allows neurons to de-couple from their current sub-population and synchronize with a different sub-population, only by adjusting the corresponding $\alpha_k$'s.

The apparent amplitude $A^*_\omega$ and phase $\phi^*_\omega$ (at frequency $\omega$) of an EEG signal are those which are obtained by performing a time-frequency decomposition of the signal. For the population model (Equation 5.1), one can obtain the apparent amplitude and phase by passing $V$ through a band-pass quadrature filter tuned at frequency $\omega$. If the filter is sufficiently narrow, the components
of $V$ with $\omega_k \neq \omega$ will be attenuated; therefore, the complex filtered signal $S$ can be approximated by:

$$S_\omega(t) = \sum_{k: \omega_k = \omega} \alpha_k e^{i(\omega t + \phi_k)},$$  \hspace{1cm} (5.2)

Since $S_\omega(t) = A_*^\omega(t) \exp[i\Phi_*^\omega(t)]$, where $\Phi_*^\omega(t) = \omega t + \phi_*^\omega(t)$ is the argument of the signal, it is possible to cancel the frequency component $e^{i\omega t}$ in the previous expressions to obtain:

$$A_*^\omega(t) \exp[i\phi_*^\omega(t)] = \sum_{k: \omega_k = \omega} \alpha_k \exp[i\phi_k].$$  \hspace{1cm} (5.3)

In other words, the apparent amplitude and phase are those of the resultant of the sum of vectors $\alpha_k \exp[i\phi_k]$, $k: \omega_k = \omega$ in the complex plane. Particularly, for two sub-populations ($k = 1, 2$, $\alpha = \alpha_1 = 1 - \alpha_2$), we have that

$$A_*^\omega)^2 = a^2 + b^2,$$

$$\tan \phi_*^\omega = b/a,$$  \hspace{1cm} (5.4)\hspace{1cm} (5.5)

with

$$a = \alpha \cos \phi_1 + (1 - \alpha) \cos \phi_2,$$

$$b = \alpha \sin \phi_1 + (1 - \alpha) \sin \phi_2.$$  \hspace{1cm} (5.6)\hspace{1cm} (5.7)

Figure 5.1 shows (for two sub-populations) how the apparent amplitude and phase vary with respect to the phase difference $(\phi_1 - \phi_2)$ and proportion $\alpha$. 

Figure 5.1: Apparent amplitude and phase versus underlying phase difference $\phi_1 - \phi_2$ and proportion $\alpha$ of oscillators with phase $\phi_1$ for a 2-population model.
Note that according to this model, a decrease of amplitude may be caused by a local resynchronization process where a group of neurons change their phase resulting in a more uniform proportion of sub-populations. Also, according to this, variations in the apparent phase may be related either to variations in the phase of some sub-populations (lack of local synchronization), or to changes in the size coefficients (which may be considered as local resynchronization processes). With this in mind, one could consider the local phase constancy (as defined by Equation 4.4) as a measure of local synchrony.

5.3 Zero-phase synchrony

With the population model one can explain why the apparent phase difference between two synchronized distant regions can be zero, in spite of the fact that there may be a significant transmission delay: with two sub-populations one can model a unidirectional coupling between two distant cortical areas, in the sense that a sub-population in one area drives a sub-population in the other area with a certain phase lag \( \delta \). For a bidirectional coupling we just add a reciprocal connection, with the same phase lag. Figure 5.2 shows a schematic of two populations in bidirectional coupling. If all sub-populations have approximately the same size (i.e. \( \alpha = 0.5 \)), then the apparent amplitude and phase for each electrode would be given by

\[
A_1^* \exp[i\phi_1^*] = 2 \cos \left( \frac{\phi_1 - \phi_2}{2} \right) \exp \left[ i \frac{\phi_1 + \phi_2}{2} \right], \quad (5.8)
\]

\[
A_2^* \exp[i\phi_2^*] = 2 \cos \left( \frac{\phi_1 - \phi_2 - 2\delta}{2} \right) \exp \left[ i \frac{\phi_1 + \phi_2}{2} \right]. \quad (5.9)
\]

So that \( \phi_1^* - \phi_2^* \approx 0 \), except in the case where

\[
\frac{\phi_1 - \phi_2 + 3\pi}{2} < \delta < \frac{\phi_1 - \phi_2 + \pi}{2}, \quad (5.10)
\]

in which case \( \phi_1^* - \phi_2^* \approx \pi \), which is in accordance with David and Friston’s neural mass model [2003].

5.4 Nodal points

In all the real EEG datasets we have analyzed, one can observe sites that show, in a specific time-frequency window, a significant increment of synchrony with a large group of electrodes. These sites act as characteristic nodes of a particular network, hence we call them nodal points. It is common, although counter-intuitive, that the group of electrodes synchronized to a nodal point show a lack of synchrony, often observed as desynchronization, among themselves. One
Figure 5.2: Bidirectional coupling between two areas with two sub-populations each. If the sub-populations are all the same size, the apparent phase difference $\phi_1^* - \phi_2^*$ will be either 0 or $\pi$.

An illustrative example of this phenomenon can be seen in the alpha band (8-12 Hz) in Figures 3.6 and 3.7, where occipital leads (electrodes T5, T6, O1, O2, and Oz) show an increase of synchrony with most of the sites, and at the same time, frontal and central leads show a synchrony decrease among them.

Let us consider three electrodes: $e_1$ shows an increase in synchrony with both $e_2$ and $e_3$. One would expect $e_2$ and $e_3$ to be also coupled, but this does not always happen. In fact, sometimes $e_2$ and $e_3$ show a decrease in synchrony. The answer, in terms of the population model, is that synchronization occurs between sub-populations (as shown in the previous section) by resynchronization processes which affect the sub-population sizes (the $\alpha_k$ coefficients), and are reflected in changes in the apparent phase which the MPD and CPPD measures can detect. A simple example of this model (for three electrodes) is shown in Figure 5.3: an increase in synchrony between $e_1$ and $e_2$ (i.e.- an increase of $\alpha_{1,1}$ and $\alpha_{2,1}$) may result in a decrease of synchrony between $e_2$ and $e_3$ since the sum of sub-population sizes must remain constant for each electrode. Of course, with a larger number of sub-populations, more complex synchronization patterns can be simulated.

On the other hand, we have observed that the relative increase in synchrony with a nodal point is usually correlated with a significant power decrease in the nodal point itself. To study this in detail, we define a site $e$ as $k$-nodal at time $t$ and frequency $\omega$ if it shows significant increase in synchronization with at least $k$ different sites. Figures 5.4a and 5.4b show, respectively, all 15-nodal sites for the Figures experiment, and the corresponding TFT map of significant power changes. In this example, one can clearly see that most of the nodal sites suffer a power decrease.

To confirm this, one can estimate the proportion of nodal sites in which
an amplitude decrease is observed: let $N_k$ be the number of $k$-nodal points in the TFT space and $A_k^-$ the number of $k$-nodal points that also show a power decrease, then the conditional probability of significant power decrease given that a site is $k$-nodal can be estimated by $P_k^- = A_k^- / N_k$. One may similarly estimate the probability of amplitude increment $P_k^+$. These probabilities are shown in Figure 5.5, where it is clear that $P_k^-$ increases with the number of couplings $k$.

This behavior may also be simulated with the population model: suppose the population covered by the nodal point is divided into various sub-populations, each one of which is coupled with a different site (see Figure 5.6a). If the sites to which the nodal point is connected are not in synchrony, the underlying sub-population phases may assumed to be fairly different. In other words, the underlying phases $\{\phi_i\}$ will show high dispersion, which will be reflected as a low apparent amplitude $A^{\omega e}$ (Equation 5.3). Figure 5.6b shows a synthetic example which simulates a nodal point placed at O2. The particular model for each signal $V_{Se}$ is given by

$$V_{Se}(t) = \sum_{k=1}^{N_k} \alpha_{e,k} \cos(\omega t + \phi_k) + \epsilon_e(t), \quad \epsilon_e(t) \sim \mathcal{N}(0, \sigma_e)$$

with $N_k = N_e$, $\omega = 10$ Hz (for this example), $\sigma_e = 0.02$, and underlying phases $\phi_k \sim \mathcal{U}(-\pi, \pi)$, $k = 1, \ldots, N_k$. In the pre-stimulus segment, neurons under site $e$ oscillate predominantly with phase $\phi_e$, thus there is little synchronous activity. This may be accomplished by assigning $\alpha_{e,e}$ a higher value than $\alpha_{e,k}, k \neq e$ for
Figure 5.4: (a) TFT map of 15-nodal points (i.e., sites which increase their synchrony with at least 15 other sites) for the Figures experiment. (b) TFT map of significant amplitude changes (red = increase, green = decrease).
Figure 5.5: Conditional probabilities of amplitude increment (red) and decrease (green) in $k$-nodal points, for the Figures experiment.

Each $e = 1, \ldots, N_e$. Specifically, we have that

$$\alpha_{e,k} = \begin{cases} 
|\eta_{e,k}|, \eta_{e,k} \sim \mathcal{N}(0, \sigma) & \text{if } k \neq e, \\
1 - \sum_{k \neq e} \alpha_{e,k} & \text{if } k = e.
\end{cases}$$

(5.12)

with $\sigma = 0.01$ for this particular example.

In the post-stimulus we model site $e_{\text{nod}}$ as a nodal point by increasing the proportion of sub-populations with phases $\phi_k, k \neq e_{\text{nod}}$ (at the expense of a lower $\alpha_{e_{\text{nod}},e_{\text{nod}}}$. This will also cause a drop in the nodal site’s amplitude, just as expected. The $\alpha_{e_{\text{nod}},k}$’s in our example are almost uniformly distributed:

$$\alpha_{e_{\text{nod}},k} = \begin{cases} 
\frac{1}{N_e} + \eta'_{e_{\text{nod}},k}, \eta'_{e_{\text{nod}},k} \sim \mathcal{N}(0, \sigma) & \text{if } k \neq e_{\text{nod}}, \\
1 - \sum_{k \neq 1} \alpha_{e_{\text{nod}},k} & \text{if } k = e_{\text{nod}}.
\end{cases}$$

(5.13)

Additionally, we have modeled a synchrony decrease among all non-nodal sites by halving the variance of the $\alpha_{e,k}$’s in Equation (5.12) (only for non-nodal sites in the post-stimulus segment). This can be seen as a resynchronization process where half the neurons of each sub-population $k$ under site $e$ change their phase from $\phi_k$ to $\phi_e$, which results in a decrease in synchrony with site $k$.

Amplitude decreases in nodal points have also been observed in three other experiments (Words and Sternberg - see Appendix ??). Two more experiments (Letters) show a different behavior: there is a high probability of observing a power increases in nodal points. In this case, however, the sites in synchrony with the nodal point show significant coupling between them, suggesting a low dispersion of the subjacent phases, which may account for the increase in amplitude.
Figure 5.6: (a) Population model applied to a nodal point: each sub-population under the nodal site is coupled with a different site (shown in blue). The blue sites show no interaction between themselves thus the phases \( \{ \phi^j \} \) may be different enough to originate a drop in the nodal point’s apparent amplitude. (b) Synchrony pattern obtained in the alpha band (post-stimulus segment) for the synthetic example of a nodal point (generated with the population model - see text for details). (c) TFT map of significant power changes from -100 to 300 ms (0 ms being the time of stimulus onset) for the synthetic nodal-point example: a clear power drop in the nodal site can be observed.
Chapter 6

Results and conclusions

6.1 Software application

The methodology described in the previous chapters has been implemented in a software program, which performs the time-frequency decomposition of the raw EEG signals, estimates any of the measures previously described, and then performs a significance test using either Bayesian classification or thresholded p-values. The results can be presented in various types of display: multitoposopes, TFT maps, and partitioned synchrony maps.

The following measures are currently implemented in the program:

- Changes in log-power with respect to the pre-stimulus (typical induced-power analysis)
- MPD
- CPPD (with threshold $\epsilon$ given in radians or ms)
- PLS
- STPLS (with window size $w$ given in ms)
- Coherence (with window size $w$ given in ms)
- LPC (with window size $w$ given in ms)
- Local phase constancy across trials (LPCT)

Additionally, the program can perform the same type of studies on evoked potentials (e.g., potentials averaged across trials), with the exception of those measures which require multiple trials to be estimated (CPPD and PLS).

For any of the synchrony measures, the program can display the following maps:
• TFT Histograms of significant synchrony increases (or decreases) for each electrode
• TF Histograms of significant synchrony increases (or decreases) across all the surface
• Multitoposcopy display of synchrony patterns for a given time and frequency
• Partitioned synchrony maps with a representative multitoposcope for each TF region
• TF maps of class labels for a given electrode pair (an example can be seen in Figure 6.6)

A full manual of the program can be found in Appendix ??.

6.2 Analysis of real EEG data

We have used the software application in collaboration with Dr. Thalía Harmony and Ph.D. student Berta González to study the dynamic synchrony changes in the Figures and Letters experiments. In this section we summarize our findings. All synchrony histograms and TFT maps for each experiment can be found in Appendix ??.

6.2.1 Results from the Figures experiment

Recall that we are measuring the changes in synchrony during a particular state in relation to a previous condition that is considered neutral. These changes can be observed, concurrently, at all frequencies. For example, there is an increase in synchrony in the delta band (1-3 Hz) in the anterior regions between 450 and 600 ms; after this time, the synchrony increased mainly in the posterior regions (Figure ??). In the first 150 ms in the delta and theta bands there was a decrease in synchrony in occipital regions (Figure ??); this may be due to the activation of the visual areas produced by the stimuli. In the theta range (4-7 Hz), the most generalized change in synchronization in the EEG after 300 ms is the phase-lock increase between posterior and anterior regions, which can be observed in the partitioned map (Figure ??). This pattern may be related to activation of Working Memory (WM), which is the process of actively maintaining a representation of information for a brief period of time so that it is available for use. Attention, decoding, perception and maintenance in memory are processes that involve the activation of WM. According to [Ungerleider et al., 1998], visual working memory involves the concerted activity of a distributed neural system, including posterior areas in visual cortex and anterior areas in prefrontal cortex.
The increase in theta (4-7 Hz) power in frontal regions (Figure ??) has been also related to activation of WM [Gevins et al., 1997; Rhom et al., 2001]. Anterior regions are also involved in encoding [Klimesch, 2004] and maintaining the information in memory [Barde and Thompson-Schill, 2002]. Sarnthein et al. [1998] report an increase in coherence in the theta range between prefrontal and posterior electrodes during retention of a string of characters.

In the alpha range (8-12 Hz), simultaneously with the increase in synchrony in the occipital regions, there was a decrease in synchrony between all regions except the occipital leads and the frontal regions on which synchrony increases. This increase may be indicating the projection loops between attentional control system in prefrontal cortex and activated meaning representations in semantic memory in posterior regions. The decrease in synchrony in the alpha band between 100 and 1500 ms coincides with the phenomena described in the 1930s by Adrian and Mathews of amplitude decrease (which can also be observed in Figure ??) or “desynchronization” and it has been related to attention [Gevins et al., 1997; Rhom et al., 2001; Klimesch, 1999].

Another change that was observed is the increase of synchrony in occipital regions in the beta band. In humans, in intracranial recordings that limited regions of extrastriate visual areas, separated by several centimeters, EEG activity in the beta range (15-25 Hz) became synchronized in an oscillatory mode during the rehearsal of an object in visual short-term memory. According to Tallon-Baudry et al. [2001] these findings confirm experimentally the hypothesis of a functional role of synchronized oscillatory activity in the coordination of distributed neural activity in humans, and support Hebb’s [1949] popular concept of short-term memory maintenance by reentrant activity within the activated network.

In the gamma band (30-40 Hz), an increase of synchrony in the interval 200-300 ms was observed. This has been related to visual search and perception [Tallon-Baudry et al., 1997]. The increase in synchronization during the interval of 600 to 800 ms around 34 Hz might be related to preparation of the motor response.

6.2.2 Results from the Letters experiment

Synchronization during the Go and NoGo conditions from the Letters experiment was studied using signals from projections to cortical sources (see Section 3.7) instead of raw potentials.

In the Go/NoGo paradigm, overall brain activity seems to involve several sub-processes, namely discrimination between the stimuli, activation of working memory, a decision whether or not to move, and initiation or inhibition of the motor actions [Shibata et al., 1999]. Anticipatory processes are also involved [Basar-Eroglu et al., 1992].

In the analysis of power changes (Figures ?? for the Go condition, and ?? for the NoGo condition) we observed an increase of delta activity from 100 to
450 ms in the Go condition and from 0-500 ms in the NoGo condition. This has been related to decision making and matching processes [Basar-Eroglu and Demiralp, 2001] and to inhibition of non-related task processes during attention to internal processes [Harmony et al., 1986]. In the Letters experiment, the delta power increase could be related to any of the processes mentioned.

Increases in theta (4-7 Hz) amplitude and synchronization have been mainly related to activation of Working Memory (WM) [Gevins et al., 1997, 1998; Rhom et al., 2001; Harmony et al., 2001, 2004]. The WM model of Baddeley [1999] distinguishes central executive functions from storage functions related to two slave systems, the phonological loop, maintaining verbal information, and a visuospatial sketch pad, responsible for the transient storage of visuospatial information. Frontal lobes have been considered the neural substrate of the central executive processes, but growing evidence has shown that executive functions do not only rely on prefrontal cortical activation but also on a distributed frontoparietal network [Sauseng et al., 2005; Osaka et al., 2004]. Visual WM involves the concerted activity of a distributed neural system which included posterior areas in the visual cortex and anterior areas in the prefrontal cortex [Ungerleider et al., 1998].

Theta activity in anterior regions is also involved in encoding [Klimesch, 1999] and in maintaining the information in memory [Barde and Thompson-Schill, 2002]. Sarnthein et al. [1998] found an increase in theta coherence between prefrontal and posterior electrodes during retention of a string of characters. These authors also reported that gamma (19-32 Hz) synchrony increased during both perception and retention intervals, suggesting that locally driven synchrony might take place in the gamma range, whereas inter-areal interactions, as in WM, appear at lower frequencies.

Shack et al. [2005] suggested that theta activity reflects central executive functions whereas upper alpha may be important for the reactivation of long-term memory codes in short-term memory.

Mazaheri and Picton [2005] reported that induced theta activity might be related to memory or attentional processes. An increase in theta power at around 300 ms after stimuli occurred with episodic memory [Klimesch, 1999]. Recognition memory, which is likely to be involved in target discrimination, has also been found to be related to synchronization of theta rhythm [Burgess and Gruzelier, 1997]. Frontal theta activity has been observed during focused attention and in association with response processing [Gevins et al., 1997]. Bastiaansen and Hagoort [2003] report that experimental evidence largely supports the hypothesis that theta activity plays a functional role in cell assembly formation, a process that may constitute the neural basis of memory formation and retrieval.

Simultaneously with the increase in power in the theta range (0-400 ms), there is also an increase of power in the alpha band in the first 200 ms, and later, a decrease in alpha power (200-1000 ms) in the Go condition. In the NoGo
condition there was a generalized increase in theta power from 0 to 500 ms, and at 8 and 9 Hz, a power increase from 0 to 450 ms. Power decrease in the Go condition was observed after 250 ms in frequencies from 8 to 29 Hz and in the NoGo condition from 600 to 900 ms in the theta range and from 50 to 300 ms at 18-20 Hz.

The decrease in power in the alpha band between 100 and 1000 ms coincides with the “desynchronization” phenomenon described in the 1930s by Adrian and Mathews, which has been related to attention [Gevins et al., 1997; Rhom et al, 2001; Klimesch, 1999]. Alpha suppression in response to effortful mental tasks has been interpreted as the marker of activation of functionally involved brain areas. However, this response is inconsistent and depends on the specific cognitive strategies employed by the subjects [Kolev et al., 2001]. In our experiment, in the Go condition, alpha power increased in the first 200 ms, and decreased afterwards. However in the NoGo condition, alpha power increased for a prolonged period up to 500 ms. Decreases in power in low alpha band in the NoGo condition were observed only from 600 to 850 ms. According to Kolev et al. [2001], enhanced early alpha oscillation to task stimuli within 0-250 ms may be related to task stimulus evaluation.

Alpha synchronization increased in both conditions (Figures ?? for the Go condition, and ?? for the NoGo condition), mainly between posterior regions and the remaining cortex. The right lateral occipital region is selectively activated during visuospatial attention [Foxe et al., 2003], which may explain our results showing this particular increase in synchronization between occipital (more in the right hemisphere) regions and the rest of the cortex.

In this experiment, one can also observe an increase of synchrony in occipital regions in the beta band. Transient long-range phase synchronization in the beta band has been related to the fronto-parietal-temporal attentional network during an attentional task [Croxson et al., 2005]. Beta synchronization across the hemispheres was reported during the perception of objects [Von Stein and Sarnthein, 1999] and during mental calculation [Mizuhara et al., 2005]. This is also supported by the report by Tallon-Baudry et al. [2001] about the role of synchronized activity in the coordination of distributed neural activity, as described in the previous section.

Oscillatory synchronization has been suggested to characterize a neuronal assembly coding for an object. Neurons distributed across different brain areas would synchronize their firing in the gamma band [Singer, 1993], supporting the idea of a role of gamma band synchrony in feature binding [Tallon Baudry et al., 1997]. In the gamma band an increase of synchrony in the first 200 ms was observed only in the Go condition. According to Karacas et al. [2001] the early high gamma response (¿75 Hz) that occurred as a phase-locked activity in the early time window of 0-150 ms is obtained regardless of stimulus type and task demands: thus, it is not a perceptual phenomenon, but instead it represents sensory processes. However, this was not confirmed with our experiments, since no gamma change was observed in the NoGo condition, although we did not
explore such high frequencies. This increase in synchrony in the gamma band
has been related to visual search and perception [Tallon-Baudry et al., 1997].
The increase in synchronization during the interval of 700 to 1300 ms in the
gamma band might be related to preparation of the motor response, since it
was observed only in the Go condition, which confirms other studies [Salenius
et al., 1996].

Multitoposcopes corresponding to a partition by bands of frequency and time
intervals gives a clear picture of the changes produced in synchrony (Figures ??
and ??, for Go and NoGo, respectively). In the Go condition, delta synchrony
increases during the whole post-stimulus interval, whereas in the NoGo condition
the synchrony increases only from 0-900 ms. In the theta band very similar
changes were observed for both conditions, with a large increase in synchrony
in the interval 300-600 ms. In the low beta band increases in synchrony were
observed in all intervals in the NoGo condition. In the high beta and in the high
gamma bands, very similar changes were observed in both conditions. However
in the low gamma band (30-40 Hz), opposite effects were observed in the two
conditions: increases in synchrony in all intervals in the Go condition but only
decreases in synchrony in the NoGo condition. These great differences may be
due to preparation and motor activation during the Go condition [Salenius et
al., 1996].

6.3 Conclusions

During the performance of different cognitive and sensorial brain tasks, several
neuronal assemblies, which are not necessarily contiguous, may become active
simultaneously and generate synchronized macroscopic oscillations. This dy-
namic reorganization produces complex patterns of synchronization and desyn-
chronization between different cortical areas, with respect to a previous state
considered as neutral. Based on this, significative increases in synchrony mea-
surements between leads indicate that the cortical areas corresponding to those
leads may be integrating a network related to a particular psychophysiological
process. Thus, EEG synchrony measures between regions may yield important
information about the dynamics of cell assemblies.

In this work, we have favored instantaneous in-phase (i.e., zero phase differ-
ence) synchrony measures, such as MPD and CPPD, for various reasons: first,
there is extensive evidence showing that, in most cases, the phase difference be-
tween two areas changes towards zero when the areas are coupled; this evidence
includes studies with real EEG data by Friston et al. [1997] and Rodriguez et
al. [1999], theoretical results from mathematical models (David and Friston’s
neural mass model, and our population model) where bidirectional couplings
result in synchrony with a phase-lag of zero or \( \pi \), reports by David and Friston
[2003b] and Varela et al. [2001] arguing that the proportion of bidirectional
connections in the brain is very high, and the high correlation between MPD,
CPPD, and PLS (which is not an in-phase measure). Second, instantaneous measures do not seem to be as affected by local phase dispersion as those measures which are estimated within a time window (e.g., coherence and STPLS). And third, our tests have shown that MPD and CPPD are not affected by volume conduction more than any of the other measures discussed here. On the other hand, coherence and STPLS may be more robust to latency jitter, and thus provide better detection of induced responses, but unfortunately, in this case, local phase dispersion may seriously mask the long-range synchronous activity.

Even if MPD and CPPD have clear advantages, other steps involved in the detection of phase synchronization must be performed with care. The first problem which concerns us is the possible spurious couplings introduced by volume conduction. Although various techniques have been proposed to reduce the volume conduction effects, none of them has proven to be sufficiently reliable. Estimation of the Surface Laplacian (SL), favored by Nunez [1995, 1997, 1999], requires a high electrode density, and thus cannot be applied to most of our datasets. On the other hand, solutions based on the inverse problem depend, in great measure, on constraints which are relatively arbitrary. In any case, one should use these solutions to complement the results obtained with raw potentials, and not as the only representation of the EEG dynamics.

Another problem is related to the time-frequency decomposition techniques used to extract the phase of the signals. As we have shown, the more popular Gabor filters and wavelets may not be adequate for low frequency analysis (delta and theta bands). In general, one must design the filters such that they have no significant response to negative frequencies. Sinusoidal quadrature filters appear to be a reliable solution to this problem.

To determine which synchrony values are significantly different from the pre-stimulus state, we subtract the baseline average, and then use a Bayesian classification method with a Markov Random Field model instead of the typical thresholded p-values. The Bayesian approach permits one to impose regularity constraints that control the granularity of the results. This is particularly important for instantaneous measures, which produce results that are relatively noisier than those measures estimated across a time window (e.g., coherence), and also has the advantage that the resulting class field is relatively robust to a wide range of values for the constraint parameter, whereas the p-value threshold, in the other case, has to be adequately chosen for each particular experiment.

Once the class label field for a particular synchrony measure has been computed, it is possible to present the results in various ways. A multitoposcopic display is useful to show the synchrony pattern corresponding to a specific time and frequency point or window. One can then plot a partitioned synchrony map by drawing a series of multitoposcopes that would cover the time segments and frequency bands of interest. On the other hand, TFT synchrony histograms can be used to display the number of connections (with respect to the baseline) in which each electrode participates: regions with high inter-connectivity may
represent integrated networks which are related to specific cognitive or sensorial processes. All these visualization tools allow the neuroscientist to explore the changes of electroencephalographic states during the course of a cognitive experiment with great detail. The analysis of the Figures and Letters experiments provide an illustrative example of the usefulness of these techniques.

Finally, a population model for the apparent phase and amplitude observed at an electrode can successfully simulate the synchrony patterns observed in real EEG data, including some interesting phenomena such as the existence and behavior of nodal points.

6.4 Future work

Here we discuss a few recent ideas that we intend to develop in the future. Additionally, and in conjunction with the UNAM Neurobiology Institute (INB-UNAM), we expect to analyze various EEG experiments, which may require specific changes in the methodology.

6.4.1 Evoked potentials and synchronization

EEG activity is typically characterized as evoked (i.e., phase-locked to the stimulus), or induced (time-locked but not necessarily phase-locked). In practice, evoked activity is revealed by averaging the EEG signals across trials, and performing a spectral analysis on the averaged signals: any activity which is not phase-locked to the stimulus will show a certain amount of phase dispersion across trials, which will result in destructive interference when averaging, leaving only the evoked responses untouched. On the other hand, induced activity is obtained by performing the spectral power analysis for each trial, and then averaging the power across trials.

One of the mechanisms which are thought to generate evoked responses is the partial phase-resetting [Makeig et al., 2002; Penny et al., 2002], in which a particular combination of the network’s current state and the network’s input cause the system’s phase to be reset to a certain value. Figure 6.1 shows the average potentials obtained from the GO condition of the Letters experiment. In this example, one can see that some of the peaks (e.g., around 150 ms and 300 ms) are observed in several sites, suggesting a relation between evoked activity and long-range in-phase and anti-phase synchronization. However, even if the phase-resetting mechanism may affect various networks located in different cortical areas, one could argue that due to the variations in the transmission delay of the reset signal, the affected areas will not necessarily be in perfect synchrony after their phase has been reset. In other words, if an evoked potential is observed in various areas, one may not necessarily observe in-phase synchronization between those areas. The opposite is also plausible: two cortical areas may increase their synchrony (with respect to the baseline), but if there is no
event-related phase-resetting mechanism affecting the subjacent phase of those areas in each trial, an evoked potential may not be generated. In any case, a measure of phase constancy across trials may yield some additional information about this type of mechanism.

Other authors (e.g., [Freeman and Schneider, 1982]) argue that evoked potentials originate, in fact, from an event-related activation of neural assemblies distinct from background dynamics [Fell et al., 2004]; in other words, the stimulus causes certain neural assemblies to generate a particular waveform, which may be reinforced by averaging across trials.

It is also possible that a mixture of both mechanisms is present in the development of evoked activity [Penny et al., 2002; Fell et al., 2004]. It would be interesting, then, to study the evoked potentials, along with other measures (e.g., induced power, synchronization, and phase constancy across trials), and try to distinguish between all possible cases. As an example of how this problem can be approached, we present in Figure 6.2 the following maps for the Letters experiment (GO condition):

(a) TFT map of changes in evoked power (EP); that is, the power of the average signals

(b) TFT map of changes in local phase constancy across trials (LPCT)

(c) SIH of evoked MPD (eMPD) synchrony (i.e., MPD between pairs of average signals)
Figure 6.2: (a) TFT map of significant EP changes (red = increase, green = decrease). (b) TFT map of significant LPCT changes (red = increase, green = decrease). (c) TFT histogram of evoked MPD increases. (d) TFT histogram of induced MPD increases. (e) TFT map of total power changes with respect to the pre-stimulus. (f) TFT map of total power minus evoked power.

(d) SIH of induced MPD (iMPD) synchrony

(e) TFT map of induced power changes with respect to the pre-stimulus

(f) TFT map of induced power minus evoked power (significance values)

Note that the first two maps (evoked power and phase constancy across trials) are very similar. This supports to some extent the theory of phase-resetting. On the other hand, there is also some correlation between evoked and induced power; in fact, the difference between induced and evoked power appears to be significant in only a few places, particularly after 350 ms (Figure 6.2f). This suggest that a certain evoked waveform develops right after the stimulus and is consistent in each trial. As time goes by, it is less likely that
event-related processes (particularly, the cognitive ones) will be phase-locked to the stimulus, and thus the differences between induced and evoked activity become more significant.

In Figure 6.1, one can see that, at around 150 ms, there is evoked activity in the alpha/low-beta range at many electrodes, which appears to be in synchrony (both in-phase and anti-phase); however, the iMPD map (Figure 6.2d) does not show such synchrony. If one closely inspects the peaks of the raw evoked potentials, one can see small differences in latency between electrodes, supporting the idea that the phase-resetting mechanisms do not affect all areas at the exact same time. It is also interesting that there is very little evoked synchrony (Figure 6.2c), compared to induced synchrony (possibly also due to the peak latency variations between leads).

Between 300 and 600 ms one can also see (in the iMPD map) a certain amount of synchronous activity in the alpha range (∼12 Hz), which does not coincide with an increase in EP.

This suggests three different cases of correlation between EP changes and induced synchrony changes between two areas, that may be of interest:

1. The two areas show an increase in evoked power and their synchronization also increases.
2. The two areas show an increase in evoked power but not in synchroniza-

3. The two areas show an increase in synchroniza-

Each of these cases may represent a different type of process. In order to obtain a detailed view of the coincidences between EP increases and induced synchrony increases, one can plot the coincidence map $C$ given by

$$C_{\omega,e_1,e_2,t} = \begin{cases} 
1 & \text{if } c_{\omega,e_1,t}^{\text{ep}} = c_{\omega,e_2,t}^{\text{ep}} = 1 \text{ and } c_{\omega,e_1,e_2,t}^{\mu} = 1 \\
2 & \text{if } c_{\omega,e_1,t}^{\text{ep}} = c_{\omega,e_2,t}^{\text{ep}} = 1 \text{ and } c_{\omega,e_1,e_2,t}^{\mu} \neq 1 \\
3 & \text{if } (c_{\omega,e_1,t}^{\text{ep}} \neq 1 \text{ or } c_{\omega,e_2,t}^{\text{ep}} \neq 1) \text{ and } c_{\omega,e_1,e_2,t}^{\mu} = 1 \\
0 & \text{otherwise}, \end{cases}$$

(6.1)

where $c^{\text{ep}}$ is the class field corresponding to EP changes, and $c^{\mu}$ is the class field corresponding to synchrony measure $\mu$. Figure 6.3 shows the coincidence map for the GO condition using MPD as synchrony measure.

Each of the three cases described above can be explained with the population model and a phase-resetting mechanism: consider the model for two bidirectionally coupled areas, and suppose that the subjacent phase of each
Figure 6.3: Partitioned map of coincidences between EP increases and iMPD increases; red means an increase in EP (at both electrodes) and an increase in iMPD ($C = 1$), green means increase in EP but not in iMPD ($C = 2$), and blue means increase in iMPD but not in EP ($C = 3$).
driving sub-population can be reset with an external signal $r$ (see Figure 6.4a). Note that $r$ is delayed at each area $i$ by $\lambda_i$, thus the subjacent phases are not necessarily reset at the exact same time. On the other hand, the coupling parameter $\alpha$ is related with the sub-population sizes; for example, perfect in-phase synchrony is obtained with $\alpha = 0.5$ (i.e.- sub-populations of the same size).

If an event-related reset signal is sent to either (or both) areas, an evoked potential (i.e.- an increase in EP) will be observed, and depending on the difference $\lambda_1 - \lambda_2$, an increase in iMPD may also been observed (case $C = 1$) even if the true synchronization between the two areas has not changed. In this case, it is not possible to distinguish if an increase in iMPD (i.e., near zero apparent phase difference) is due to a true increase in synchronization, or due to a phase-resetting mechanism which acts upon both areas almost at the same time. On the other hand, if $\lambda_1 - \lambda_2$ is large enough, resetting the subjacent phases in both driving sub-populations will not result in spurious high synchrony (case $C = 2$).

If there is phase-resetting but no true coupling (Figure 6.4b), one will observe an increase in EP, and may or may not observe a (spurious) increase in iMPD (cases $C = 1$ and $C = 2$, respectively).

The last case ($C = 3$) is explained, according to the model in Figure 6.4c, where there is coupling but no phase-resetting. The change in iMPD may be caused simply by a change in the subjacent sub-population sizes (i.e.- a resynchronization process where more neurons become part of the sub-populations involved in the bidirectional coupling), such that $\alpha$ changes towards 0.5. The idea of neurons resynchronizing to a different sub-population is supported by studies by Haalman and Vaadia [1998], which show that when performing a computational task, neurons may quickly associate into a functional group while disassociating from concurrently activated groups.

David et al. [2006] have also studied the mechanisms which generate evoked and induced responses (in terms of power), from the viewpoint of their neural mass model [David et al., 2003]. According to them, there are two types of mechanisms: dynamic mechanisms, which consist of modulations of the neuronal input, and structural mechanisms, which correspond to changes in the system’s causal structure (e.g., the model parameters). David et al. study how a mixture of these mechanisms is reflected as evoked and induced responses.

In our population model, the resetting function $r$ is the system’s input, thus one can consider the phase-resetting mechanism as a dynamic one. Furthermore, the resynchronization of neurons from one sub-population to another may be considered a structural mechanism. Our intention is to study in more depth these mechanisms, and their expression in terms of evoked power, induced power, and long-range phase synchronization.

6.4.2 Anti-phase synchrony

Earlier, we discussed the concept of anti-phase synchronization; that is, coupling with a phase difference of $\pi$ radians. Both, David and Friston’s neural mass
Figure 6.4: Population model for two bidirectionally coupled areas (with coupling parameter $\alpha$), with a phase-resetting signal $r$ affecting the driving sub-populations.
model, and our population model, yield a phase difference of zero or π for two areas which are bidirectionally coupled. Moreover, as can be seen in Figure 6.1 (around 150 ms), when some electrodes show a positive evoked potential, others display a negative peak at approximately the same latency, suggesting the presence of anti-phase synchrony.

Therefore, it would be interesting to develop a method to detect true anti-phase couplings, and study its relation with in-phase synchrony and evoked potentials. As a first approach, an anti-phase CPPD (AP-CPPD) measure has already been proposed in Section 3.3.3 (Equation 3.7). The problem with this measure is that it is highly correlated (negatively) with the in-phase CPPD. This means that it is not easy to distinguish, with these measures, between increases in anti-phase synchrony and decreases in in-phase synchrony. One thing that helps is using small values for the threshold $\epsilon$, at the expense of a more conservative criteria for synchrony (e.g., fewer significative changes with respect to the baseline).

An example of these measures is shown in Figure 6.5. Here we present the synchrony decrease histograms of the (in-phase) CPPD measure, and synchrony increase histograms of the anti-phase CPPD, for $\epsilon$ values of $\pi/5$ and $\pi/20$. 
For larger $\epsilon$ values, the CPPD SDH and AP-CPPD SIH are relatively similar, whereas smaller thresholds seem to be more adequate to discern between the anti-phase synchronization and desynchronization.

### 6.4.3 Synchrony at multiple frequencies

Another interesting phenomenon is the ability of two electrodes to display in-phase synchronization at various frequencies, at the same time. To see this, one can plot the TF map of class labels for a particular pair of electrodes. Figure 6.6 shows a few examples of this behavior for pairs Fp1-Fp2, Fp1-O2, and T5-T6 of the Figures experiment. The case of T5-T6 is particularly interesting since there is an increase in synchronization in the delta and theta bands, a decrease in the alpha and low beta bands, and then another increase in the beta and gamma bands. This may be explained by the idea (already discussed in Section 5.1) that during certain tasks, neurons may become part of a functional group, while disassociating from other groups. If these functional groups generate macroscopic oscillations at different frequencies, then one may be able to see synchronization patterns which are multiplexed across a wide frequency range, such as those in Figure 6.6. It is possible, then, that in order to obtain a full picture of the synchronization pattern at a given time, one should integrate the information for all frequency bands in a single display.
Bibliography


