Estandarización y automatización en el diagnóstico de pacientes con desórdenes de conciencia: aprendizaje automático aplicado a señales electrofisiológicas del cerebro y el cuerpo

Raimondo, Federico
2018

Este documento forma parte de la colección de tesis doctorales y de maestría de la Biblioteca Central Dr. Luis Federico Leloir, disponible en digital.bl.fcen.uba.ar. Su utilización debe ser acompañada por la cita bibliográfica con reconocimiento de la fuente.

This document is part of the doctoral theses collection of the Central Library Dr. Luis Federico Leloir, available in digital.bl.fcen.uba.ar. It should be used accompanied by the corresponding citation acknowledging the source.

Cita tipo APA:

Cita tipo Chicago:
Estandarización y automatización en el diagnóstico de pacientes con desordenes de conciencia: aprendizaje automático aplicado a señales electrofisiológicas del cerebro y el cuerpo.

Tesis presentada para optar al título de Doctor de la Universidad de Buenos Aires en el área Ciencias de la Computación

Federico Raimondo

Directores de tesis: Dr. Diego Fernández Slezak
                     Dr. Jacobo Diego Sitt
                     Dr. Laurent Cohen

Consejero de estudios: Dr. Agustín Gravano

Buenos Aires, Septiembre de 2018
Standardisation and automatisation of the diagnosis of patients with disorders of consciousness: a machine learning approach applied to electrophysiological brain and body signals.

Thesis presented to apply for the degree of Doctor of the University of Buenos Aires in the area of Computer Science

Federico Raimondo

Advisors: Dr. Diego Fernández Slezak
Dr. Jacobo Diego Sitt
Dr. Laurent Cohen
Study Counselor: Dr. Agustín Gravano

Buenos Aires, Septiembre 2018
Contents

Resumen vii

Summary ix

Acknowledgments xi

1 Introduction 1

1.1 The study of Consciousness . . . . . . . . . . . . . . . . . . . . . . . . 2

1.1.1 Conscious Access vs Conscious States . . . . . . . . . . . . . . . . . 2

1.1.2 Disorders of Consciousness . . . . . . . . . . . . . . . . . . . . . . . . 2

1.1.3 Why diagnosis is important? . . . . . . . . . . . . . . . . . . . . . . . 3

1.2 State of the art of DOC diagnosis . . . . . . . . . . . . . . . . . . . . . . 5

1.2.1 Current tools for DOC diagnosis . . . . . . . . . . . . . . . . . . . . . 5

1.3 Electroencephalography . . . . . . . . . . . . . . . . . . . . . . . . . . . . 9

1.3.1 What is EEG? . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 9

1.3.2 What do we analyse in EEG? . . . . . . . . . . . . . . . . . . . . . . 11

1.4 Electroencephalography in DOC patients . . . . . . . . . . . . . . . . . . 14

1.5 Methods used in this thesis . . . . . . . . . . . . . . . . . . . . . . . . . . 17

1.5.1 The Coma Recovery Scale (Revised) . . . . . . . . . . . . . . . . . . 17

1.5.2 The Local-Global paradigm . . . . . . . . . . . . . . . . . . . . . . . . 17

1.5.3 Machine Learning . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 18

1.5.4 Statistics . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 21

1.6 This work . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 21

1.6.1 Purpose . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 21

1.6.2 Description of chapters . . . . . . . . . . . . . . . . . . . . . . . . . . 22

2 Automation and identification of robust EEG-extracted markers for the diagnosis of DOC 27

2.1 Background . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 28

2.2 Objectives . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 29
CONTENTS

2.3 Automation ................................................................. 29
  2.3.1 Reasons to rewrite the code and automate the process .... 29
  2.3.2 Software Implementation ........................................ 30
  2.3.3 Preprocessing ...................................................... 30
  2.3.4 Optimisations and Performance .................................. 31
2.4 Replication ............................................................... 31
  2.4.1 Methods ............................................................. 34
  2.4.2 Results ............................................................. 37
2.5 Validation ................................................................. 37
  2.5.1 Methods ............................................................. 37
  2.5.2 Results ............................................................. 38
2.6 Clinical application .................................................... 38
  2.6.1 Report ............................................................... 38
2.7 Further steps: towards a universal tool .............................. 40
  2.7.1 Generalisation to different EEG recording configurations ... 42
  2.7.2 Generalisation to Resting State .................................. 43
  2.7.3 Discussion .......................................................... 45
2.8 Insights from Machine Learning ...................................... 48
  2.8.1 Methods ............................................................. 48
  2.8.2 Results ............................................................. 49
  2.8.3 Discussion .......................................................... 49
2.9 Summary ................................................................. 51

3 Towards cross-modal integration as a measure of consciousness 55
  3.1 Background ............................................................ 56
  3.2 Hypotheses ............................................................. 58
  3.3 Methods ............................................................... 59
    3.3.1 Cross-modal Local-Global ...................................... 59
    3.3.2 Subjects .......................................................... 61
    3.3.3 EEG acquisition ................................................ 62
    3.3.4 Analysis .......................................................... 62
    3.3.5 Group-Level analysis .......................................... 63
    3.3.6 Statistics ........................................................ 63
  3.4 Results ............................................................... 63
    3.4.1 ERP analysis .................................................... 63
    3.4.2 ERP characterisation through MVPA ........................... 64
  3.5 Discussion ............................................................ 64
4  Brain-Body interactions as a diagnostic marker for DOC  73

4.1  Background  ................................................................. 74

4.2  Hypotheses ................................................................. 75

4.3  Baseline cardiac activity ............................................... 75

  4.3.1  Methods ................................................................. 75

  4.3.2  Results ................................................................. 77

4.4  Cardiac markers of cognitive processing  ......................... 83

  4.4.1  Methods ................................................................. 83

  4.4.2  Results ................................................................. 85

4.5  EEG and EKG Multivariate Pattern Analysis ....................... 85

  4.5.1  Methods ................................................................. 85

  4.5.2  Results ................................................................. 87

4.6  Discussion ................................................................. 89

5  Summary and final remarks  97
**Estandarización y automatización en el diagnóstico de pacientes con desordenes de conciencia: aprendizaje automático aplicado a señales electrofisiológicas del cerebro y el cuerpo.**

Avances en la medicina moderna han llevado a un incremento en el número de pacientes diagnosticados con desordenes de consciencia (DOC). En estas condiciones, los pacientes se encuentran despiertos, pero no muestran signos de entendimiento acerca de sí mismos o el lugar donde se encuentran. Una evaluación precisa de los pacientes tiene implicaciones medico-éticas y sociales, y es de suma importancia porque típicamente informa el pronóstico. Los diagnósticos erróneos, no obstante, es una gran preocupación en las clínicas debido a las limitaciones intrínsecas de las herramientas de diagnóstico basados en comportamiento. Una tecnología accesible para asistir a los médicos es la electroencefalografía (EEG). En un estudio previo, introducimos el uso de marcadores extraídos de EEG en combinación con aprendizaje automático como una herramienta para el diagnóstico de pacientes DOC. En este trabajo, desarrollamos una herramienta de análisis automatizado, y analizamos la aplicabilidad y limitaciones de este método. Adicionalmente, proponemos dos enfoques para incrementar la precisión del diagnóstico: (1) el uso de múltiples modalidades de estimulación para incluir los correlatos neuronales de la integración multisensorial y (2) el análisis de las modulaciones de la actividad cardíaca mediadas por la conciencia. Nuestros resultados exceden el conocimiento actual en dos dimensiones. Clínicamente, encontramos que el método puede ser utilizada en contextos heterogéneos, confirmando la utilidad del aprendizaje automático como una herramienta para el diagnóstico clínico. Científicamente, nuestros resultados resaltan que las interacciones entre el cerebro y el cuerpo pueden ser el mecanismo fundamental para sostener la fusión de múltiples sentidos en una única percepción, conduciendo a la emergencia de la consciencia. En conjunto, este trabajo ilustra la importancia del aprendizaje automático para la evaluación clínica individualizada, y crea un punto de partida para la inclusión de las funciones corporales en la cuantificación de los estados de conciencia globales.
Standardisation and automatisation of the diagnosis of patients with disorders of consciousness: a machine learning approach applied to electrophysiological brain and body signals.

Advances in modern medicine have led to an increase of patients diagnosed with disorders of consciousness (DOC). In these conditions, patients are awake, but without behavioural signs of awareness. An accurate evaluation of DOC patients has medico-ethical and societal implications, and it is of crucial importance because it typically informs prognosis. Misdiagnosis of patients, however, is a major concern in clinics due to intrinsic limitations of behavioural tools. One accessible assisting methodology for clinicians is electroencephalography (EEG). In a previous study, we introduced the use of EEG-extracted markers and machine learning as a tool for the diagnosis of DOC patients. In this work, we developed an automated analysis tool, and analysed the applicability and limitations of this method. Additionally, we proposed two approaches to enhance the accuracy of this method: (1) the use of multiple stimulation modalities to include neural correlates of multisensory integration and (2) the analysis of consciousness-mediated modulations of cardiac activity. Our results exceed the current state of knowledge in two dimensions. Clinically, we found that the method can be used in heterogeneous contexts, confirming the utility of machine learning as an automated tool for clinical diagnosis. Scientifically, our results highlight that brain-body interactions might be the fundamental mechanism to support the fusion of multiple senses into a unique percept, leading to the emergence of consciousness. Taken together, this work illustrates the importance of machine learning to individualised clinical assessment, and paves the way for inclusion of bodily functions when quantifying global states of consciousness.
Acknowledgments

A pesar de que es encuentra al principio, esta es la última página que he escrito sobre este trabajo. Vengo pensando en este texto desde el momento que abrí el primer archivo y comencé a escribir. Han pasado poco más de cinco años, tres países, cinco veranos, siete inviernos, tres laboratorios, y en cada uno de esos momentos, incontables personas que han formado parte de este presente en el que me encuentro. Es mi deseo agradecer a todas esas personas, a las que fueron y a las que son parte de mi vida.

Quiero agradecer a mis padres, Martha y Dicky, que desde chico me empujaron, pero me dejaron libre para elegir mi futuro (aunque nunca me dieron la posibilidad de vivir del rock).

A mis hermanas y hermano, Nina, María y Tomás, quienes tuvieron que soportar mudanzas, trámites, pedidos y favores. Quienes estuvieron siempre, en las buenas, en las malas, aun cuando yo no estuve presente.

A mis tíos, tías, abuelos y primos: Magdalena, Pepe, Diana, Leticia, Julia, Tadeo, Candela. Que en cada visita a la Patagonia me recordaban que la familia sigue estando ahí. A mi tío Gustavo, quién fue una gran influencia en mi infancia y adolescencia, y que me sigue demostrando que hay que seguir luchando a pesar de las desgracias y malos momentos. ¡Fuerza!

A mis amigos patagónicos, que a pesar de la distancia, siempre fueron un lugar de confort y alegría. En orden alfabético y sin artículos: Adriano, Chacha, Chiqui, Chomba, Churi, Eze, Foco, Gato, Guille, Herni, Juanpa, Pablito, Pantro, Rata, Rodri, Ruso, Topo. Sin olvidar Córdoba y Rosario.

A Tut, imperceptible y presente al mismo tiempo.
A Yorgos, por las noches de videojuegos para despejar la mente.
A la Abuela Lidia, quien en su tiempo proveyó de un lugar de encuentro e intercambio entre amigos. Pero más que nada, a esos amigos, que religiosamente asistían a discutir nuevas formas de mejorar el mundo entre vinos y bondiolas.

Al Departamento de Computación, por la excelente educación recibida y por ser una comunidad que no afloja, que te forma como estudiante, docente y persona.

Al PICNIC Lab, un equipo con múltiples formaciones, pero siempre un equipo, con jugadores de primera como Mariam, Diana, Sami, Denis, Camille, Florence, Benjamin,
Melanie, Pauline, Bertrand, Kasia, Tal y todos los que pasaron mientras estuve ahí y me mostraron que la ciencia excelente también puede ser divertida.

Al Coma Science Group, el lugar donde escribo esto, por darme la oportunidad de seguir mi carrera.

A Diego, que un día me encontró en un pasillo y me propuso un proyecto que arrancó con mi carrera.

A Lionel, por su confianza.

A Jaco, que se puso la cinta, agarró el timón, y empezó a remar solo en un bote que iba a la deriva, llevando este doctorado a culminar, y de una manera en la que yo nunca creí posible. Quien me formó como neurocientífico, quien me dio dirección, quien confió en mí y que, principalmente, me enseñó cómo hacer ciencia de la buena.

Y por último, pero sin quitar importancia, a esos dos hermosos seres humanos con los que comparto mi vida todos los días, que soportaron todo, los buenos y los malos momentos, y que no paran de darme energía. A Athena y Alex.

Durante el desarrollo de este trabajo, recibí financiamiento del Departamento de Computación (FCEyN, UBA), ICM (Francia) y, en particular, una beca doctoral de CONICET, que me sostuvo durante 5 años. Por este motivo, quiero agradecer especialmente a todos y todas/as que lucharon y siguen luchando por la ciencia argentina.
Chapter 1

Introduction

Contents

1.1 The study of Consciousness ........................................ 2
  1.1.1 Conscious Access vs Conscious States ....................... 2
  1.1.2 Disorders of Consciousness ................................. 2
  1.1.3 Why diagnosis is important? ............................. 3

1.2 State of the art of DOC diagnosis ............................... 5
  1.2.1 Current tools for DOC diagnosis ............................ 5

1.3 Electroencephalography .......................................... 9
  1.3.1 What is EEG? ............................................. 9
  1.3.2 What do we analyse in EEG? .............................. 11

1.4 Electroencephalography in DOC patients ......................... 14

1.5 Methods used in this thesis .................................... 17
  1.5.1 The Coma Recovery Scale (Revised) ....................... 17
  1.5.2 The Local-Global paradigm ............................... 17
  1.5.3 Machine Learning ....................................... 18
  1.5.4 Statistics ............................................. 21

1.6 This work .................................................. 21
  1.6.1 Purpose ............................................... 21
  1.6.2 Description of chapters .................................. 22
CHAPTER 1. INTRODUCTION

1.1 The study of Consciousness

1.1.1 Conscious Access vs Conscious States

Throughout history, the term consciousness has been studied from disciplines as philosophy, psychology and biomedical sciences. Nowadays, there is still no universal definition for such an ambiguous concept (Baars, 2015). Depending on the use of the word conscious, it can refer to the state of consciousness or to conscious access or processing. In its intransitive use (e.g. “the patient is still conscious”), it refers to the state of consciousness, also called wakefulness or vigilance. In its transitive use (e.g. “I was not conscious of the red light”), it refers to access or processing of a specific piece of information. The information accessed consciously at a given moment defines the conscious content, which can be reported (Dehaene and Changeux, 2011).

1.1.2 Disorders of Consciousness

The state of consciousness, at the same time, is a multidimensional construct (Bayne et al., 2016). This means that it can be expressed in several distinct states which describe arousal level, cognitive function, and bodily states. In lack of a full description of these dimensions of each state of consciousness, we will here adopt a simplified definition coming from clinical practice (Posner et al., 2007). Clinical neurologists typically evaluate consciousness on two dimensions, arousal and awareness. Arousal refers to the level of wakefulness and is clinically indicated by eyes-opening. Awareness refers to the contents of consciousness and it is clinically evaluated by command following and by observing nonreflex behaviours. What is further known from clinics is that the relationship between these two dimensions is not always a positive one. The introduction of the mechanical ventilator in the 1950s and the development of intensive care in the 1960s made it possible for many patients to sustain their vegetative functions and allowed them to survive their severe injuries. Despite such advancement, many patients were found to suffer from altered states of consciousness, which had never been encountered before as these patients would normally die from apnea (Laureys and Boly, 2007). Patients in a vegetative state/unresponsive wakefulness syndrome (VS/UWS), although they show intermittent periods of wakefulness, they do not respond to stimulation evidencing awareness of self and the environment (Jennett and Plum, 1972). Even when patients show discernible signs of behavioural non-reflex activity, coined as the minimally conscious state (MCS), they remain unable to communicate and are considered with impaired consciousness (Giacino et al., 2002). When patients regain the capacity of functional communication or object user are considered to be emerged from MCS (EMCS). An illustration comparing the different consciousness states is shown in...
1.1. THE STUDY OF CONSCIOUSNESS

Figure 1.1 on the following page.

1.1.3 Why diagnosis is important?

The impact of such profound states of unconsciousness is reflected in the composition of the first bioethical committees discussing the redefinition of life and death, hence predicting the medico-ethical legal, and societal debates that were to follow (Beecher et al., 1968). Debates of this kind mainly stem from how consciousness is considered in these conditions (Racine et al., 2008). For example, with a wide European survey among healthcare professionals there was a unanimous support for pain perception in MCS (96%) but less for the VS/UWS (56%) (Demertzi et al., 2009). Similarly, the majority (66%) of healthcare professionals agreed to withdraw life-sustaining treatment from chronic VS/UWS patients whereas only 28% agreed so for the chronic MCS (Demertzi et al., 2011). Additionally, disorders of consciousness have required the mediation of legal authorities in order to regulate end-of-life decisions (Quill, 2005): in the absence of a written statement about end-of-life preference from patients’ behalf (advance directive), a surrogate decision maker is eligible to mediate trying to maximise patients’ self-determination and protect their interests (Bernat, 2002, 2004). As such, conflicts of interest among caregivers can arise leading to wide societal debates (Quill, 2005; Striano et al., 2009). Also, treatment resources are not unlimited. The allocation of medical resources and the economics at the end of life have not yet been fully determined for patients for whom the dilemma on treating becomes crucial either because treatments are not guaranteed as successful (i.e., the condition is too bad to be treated) or unkind (i.e., the quality of life of those surviving is not acceptable) (Fins, 2003). Finally, the ethical significance of consciousness has raised many discussions as to whether greater sentence entails greater quality of life. As being conscious entails being conscious both of wellbeing and suffering (Kahane and Savulescu, 2009), it might hence not be in patients’ best interest to preserve life-sustaining aids (Horne, 2009).

Taken collectively, to evaluate consciousness in noncommunicating patients has medico-ethical significance. To date, efforts are focused on determining reliable diagnostic labels. Diagnosis is considered a crucial level, because it typically informs about patients’ prognosis, i.e., unresponsive patients have less favourable outcome as compared to those in MCS (Faugeras et al., 2017), and may influence clinical management and treatment options (Jox et al., 2012).
CHAPTER 1. INTRODUCTION

Content of consciousness (awareness)
Level of consciousness (wakefulness)
Conscious
Wakefulness
Drowsiness
Sleep stages I-II
Deep Sleep

“Vegetative” / Unresponsive Wakeful Syndrome
Minimally Conscious State
MCS + (command following)
MCS - (non-reflex movement)
General Anaesthesia
Coma

Figure 1.1: Simplified illustration of the two major components of consciousness: the level (wakefulness) and the content of consciousness (awareness). Normal physiological states (purple) present a positive correlation. Patients in coma or anaesthesia (gray) are unconscious and they cannot be awakened. DOC patients are awake but do not present signs of awareness (VS/UWS patients; blue) or show inconsistent but discernible signs of behavioural activity (MCS patients; green). Adapted from Laureys 2005b
1.2. STATE OF THE ART OF DOC DIAGNOSIS

1.2 State of the art of DOC diagnosis

1.2.1 Current tools for DOC diagnosis

Behavioural evaluation

Voluntary and reflexive behaviours are difficult to distinguish. Clinicians and caregivers could miss the subtle signs of consciousness behaviour present in MCS patients. Despite the specification of a diagnostic criteria for MCS (Giacino et al., 2002), a study showed that non-standardised observation presented a 41% misdiagnosis of VS/UWS patients when compared with a standardised scale (Schnakers et al., 2009). Among the several sources of variance that contribute to this misdiagnosis, the patient’s fluctuations in terms of arousal, fatigue, illness, pain, cortical sensory deficits, motor impairments or cognitive disturbance decreases the probability of observing signs of consciousness (Schnakers et al., 2015). Taking this fact into account, behavioural evaluations should occur repeatedly over time and should be sensitive enough to detect this subtle but meaningful fluctuations.

Conventional bedside assessment procedures as the Glasgow Coma Scale (GCS; Teasdale and Jennett 1974) are designed to detect gross changes in behaviour rather than reflexive from voluntary behaviour. An alternative scale which present higher sensitivity in detecting levels of brainstem function in the acute stage of brain injury is the Full Outline of UnResponsiveness score (FOUR score) (Wijdicks et al., 2005). Nevertheless, this score does not include a systematic assessment of signs of consciousness (Giacino et al., 2002) and it may not capture the transition from VS to MCS (Schnakers et al., 2006; Bruno et al., 2011).

Standardised neurobehavioral assessment measures tailored for DOC patients include the Coma Recovery Scale – Revised (CRS-R) (Giacino et al., 2004), the Coma-Near Coma Scale (CNC) (Rappaport et al., 1992), the Western Neurosensory Stimulation Profile (WNSSP) (Ansell and Keenan, 1989), the Western Head Injury Matrix (WHIM) (Shiel et al., 2000), and the Sensory Modality Assessment and Rehabilitation Technique (SMART) (Wilson and Gill-Thwaites, 2000). Although item content varies across measures, all evaluate behavioural responses to a variety of auditory, visual, motor, and communication prompts.

Neuroimaging

Nowadays, behavioural assessment of DOC remains the ‘gold standard’ due to the lack of an objective test of consciousness. Nevertheless, neuroimaging permits objective documentation of central nervous system damage after acquired brain injury. Scientifically, neuroimaging studies contributes to a better understanding of the neural correlates of
human consciousness. Clinically, they provide additional information concerning diagnosis, prognosis and the course of recovery of consciousness, and can serve as surrogate markers for novel therapeutic interventions (Giacino et al., 2014).

Structural neuroimaging techniques used in DOC comprises Magnetic Resonance Imaging (MRI) and Computed Tomography (CT). These images allow the visualisation of the location and extent of brain damage, but cannot reliably differentiate VS/UWS from MCS patients. A previous study showed that structural MRI can predict the outcome of DOC patients; for example, the presence of corpus callosum and dorsolateral brain-stem lesions correlates with the lack of recovery at the group level (Kampfl et al., 1998). However, the recently developed quantitative diffusion tensor imaging (DTI) techniques, which permit assessment of structural white matter damage, have been shown to outperform clinical markers in predicting 1-year functional outcome at the individual-patient level in patients with traumatic (Galanaud et al., 2012) or anoxic (Luyt et al., 2012) brain injury.

Functional neuroimaging can measure brain function, either in active or passive (resting) states. Among the available technologies, functional MRI (fMRI), F-fluorodeoxyglucose PET (FDG-PET) and Electroencephalography (EEG) have helped understanding DOC. Each technique measures different aspects: FDG-PET measures the brain’s metabolic activity, fMRI the haemodynamic activity and EEG the electrical activity.

A recent study used FDG-PET to measure whole-brain glucose metabolic state, concluding that this quantification allows accurate diagnosis and prediction of disorders of consciousness (Figure 1.2 on the next page; Stender et al. 2014). Nevertheless, in the same work, the authors depict the need for high standards in image quality and registration.

Using fMRI, studies have been carried using auditory, tactile or visual stimuli, depicting near-normal cortical activation patterns in MCS patients contrary to the low-level activations in VS/UWS patients (Laureys and Schiff, 2012). Despite their potential value as prognostic markers, the diagnostic value and interpretation of activation fMRI studies in DOC in terms of the presence or absence of residual consciousness have remained controversial. Indeed, in the absence of a full understanding of the neural correlates of consciousness, deficient cortical activation to external stimuli does not necessarily prove the absence of consciousness (Laureys, 2005a). On the other hand, task-free resting-state fMRI measurements are also used for diagnosis of DOC. In a recent study, automatic classification using functional connectivity between left auditory, right auditory and occipital cortex was able to determine the conscious state of 20 out of 22 patients (figure 1.3 on page 8; Demertzi et al. 2015).
Figure 1.2: FDG-PET images for representative group samples as extracted from Stender et al. 2016.
Figure 1.3: Regions showing higher functional connectivity in MCS patients as compared to VS/UWS patients (extracted from Demertzi et al. 2015).
Electrophysiology
Another technique consists of studying the electrical activity of neurons or muscles. The term electrophysiology encompass several particular techniques, depending on the source of the signals. A common technique used throughout this entire thesis is electroencephalography (EEG) which focus on measuring the electrical activity of the cerebral cortex.

1.3 Electroencephalography

1.3.1 What is EEG?

How does it work?

Electroencephalography (EEG) is an electrophysiological technique for the recording of electrical activity arising from the human brain. The history of EEG goes back to the 19th century, starting by Richard Caton in 1875 who performed the first known neurophysiologic recording of animals using a galvanometer. It was not until 1924 when the first studies in human began, made by Has Berger, a German psychiatrist (St. Louis and Frey, 2016).

EEG uses the principle of differential amplification, or recording voltage differences between different points using a pair of electrodes that compares one active exploring electrode site with another neighbouring or distant reference electrode. This differences are measured in volts (typically microvolts or $\mu V$).

An EEG acquisition system consists on a set of delicate electrodes connected to a set of differential amplifiers (one for each channel), following by filters. Before the digital era, this amplified signals were written by a mechanical pen directly into a rolling paper, leaving ink traces of waveforms that mirrored the electrical potential differences on the scalp. Modern systems contains analog-to-digital converters (ADC) that transforms the analog signal into discrete digital data, readable by a computer or microprocessor which can be used to store the acquired signals (Sanei and Chambers, 2007).

The electrodes used to sense the voltage differences are crucial for acquiring high quality data. Nowadays, electrodes can be used to record neuronal activity in the scalp (EEG), brain cortex (Electrocorticography; ECoG), brain tissue (Local Field Potentials; LFP) or single neurons (single-unit recordings). Nevertheless, except for scalp EEG, the mentioned techniques are invasive and requires surgical procedures. Scalp EEG electrodes are pasted or glued to the scalp. For multichannel recordings with a large number of electrodes, caps are often used.
What do we measure?

The nerve cells in the brain produce signals that are called action potentials. These action potentials move from one cell to another across a gap called the synapse. Special chemicals called neurotransmitters help the signals to move across the gap. There are two types of neurotransmitters, one will help the action potential to move to the next cell (excitatory), the other will stop it moving to another nerve cell (inhibitory).

The electrical activity measured by scalp EEG recordings is generated by similarly oriented groups of cerebral cortical neurons near the scalp where the recording electrodes are placed. Each scalp electrode collects, at a minimum, an estimated 6 cm² synchronous cortical activity. The majority of the electrical activity collected in the EEG is generated by groups of pyramidal neurons. These cells have cell bodies primarily in layers three and five of the cerebral cortex. The electrical activity recorded on the scalp represents the summation of the inhibitory or excitatory postsynaptic potentials (not action potentials, they are too short to be recordable) from thousands of pyramidal cells near each recording electrode. This summated activity can be represented as a field with positive and negative poles (dipole).

There are systematic interconnections between cortical neurons, as well as cortical to subcortical connections to structures such as the thalamus, that have well-developed feed-back linkages. Any sinusoidal rhythmic activity seen on the EEG is thought to represent oscillatory communications between the cortex and deeper, subcortical structures. These communication loops occur when the cortex is at rest or is not performing any specific task. Once the cortex has a task to perform, the electrical activity of the cortex desynchronises, and lower amplitude, faster electrical rhythms take predominance until the cortex completes its task and returns to a resting state (St. Louis and Frey, 2016).

Resolution, precision and accuracy of EEG

The differences between precision, resolution and accuracy are subtle but important. Resolution refers to the number of data samples by unit time, precision refers to the certainty of the measurements at each time point and accuracy to the relationship between the timing of the EEG signal and the biophysical event that lead to that signal.

The temporal resolution of the EEG is given by the sampling rate of the acquisition, generally between 100 Hz and 20 Khz, depending on the purpose of the acquisition. For most analysis on scalp EEG, resolutions between 250 Hz and 1000 Hz are sufficient and appropriate. In contrast, the temporal precision depends on the analysis applied. Raw (unprocessed data) have the highest temporal precision because each sample was taken at a precise time point. Nevertheless, filtered data is a weighted average of the temporal
surrounding activity, which reduces the temporal precision. In terms of accuracy, the EEG is extremely accurate because brain electrical activity travels instantaneously from the neurons generating the electrical field to the electrodes measuring them.

Although EEG has high temporal precision, resolution and accuracy, it is not the case for spatial properties, as they are considered low compared to neuroimaging techniques such as fMRI. The spatial resolution of EEG is determined by the number of electrodes. Common configurations consist of 21 electrodes (10-20 standard; Jasper 1958), 32, 64, 128 and 256. The spatial precision of the EEG is considered low, although it can be improved by spatial filters. In terms of accuracy, the problem is that one electrode does not reflect the activity from neurons directly below that electrode, but rather a complex mixture of activities from many brain regions close to and distant from it. Furthermore, the extent to which one brain region contributes to the signal recorded from each EEG electrode depends on cortical anatomy and to what extent that brain region is active at a given point in time (Cohen, 2014).

### 1.3.2 What do we analyse in EEG?

#### Oscillations

An oscillation is a rhythmic alternation of states. They can occur in time or in space, and are commonly seen in physical and biological systems. In the brain, they refer to fluctuations in the excitability of neurons or populations of neurons. Neural oscillations are observed on many spatial and temporal scales (Varela et al., 2001) and have been linked to many neurobiological events ranging from long-term potentiation to conscious perception (Buzsáki, 2009; Engel et al., 2001; Herrmann et al., 2010; Kistler et al., 2000; Klimesch et al., 2007; McBain and Kauer, 2009). These neurobiological mechanisms are fairly well understood (Buzsáki et al., 2012; Wang, 2010) although uncertainties remain in the extent to which different factors contribute to the signal recorded by EEG, in part due to the complexity of the models and the difference in spatial scale between individual neurons and scalp EEG.

Brain rhythmic activity contains multiple frequencies simultaneously, which can be separated through signal processing techniques. These rhythms are grouped into bands, defined as delta (2-4 Hz), theta (4-8 Hz), alpha (8-12 Hz), beta (15-30 Hz), lower gamma (30-80 Hz), upper gamma (80-150 Hz). Although there are more oscillations, these bands are most typically associated with cognitive processes, defined from neurobiological mechanisms of brain oscillations, including synaptic decay and signal transmission dynamics (Buzsáki, 2009; Buzsáki and Draguhn, 2004; Kopell et al., 2010; Niedermeyer and Silva, 2004; Wang, 2010).

Changes in rhythmic activity correlate with task demands, including perceptual,
cognitive, motor, linguistic, social, emotional, mnemonic and other functional processes (Cohen, 2014).

Event-related Potentials (ERPs)

One way to study the human brain function is by analysing the reactions of the brain to a variety of stimuli. Some of these reactions may be associated with clear cut changes in the EEG; some others, however, consist of changes that are difficult to visualise. These EEG changes may be treated globally under the common term event-related potentials (ERPs); a subset of the ERPs are sensory (visual, auditory, somatosensory) evoked potentials (ERPs).

ERPs are usually defined in the time domain as the brain electrical activity that is triggered by the occurrence of particular events or stimuli. A basic problem of analysis is how to detect ERP activity within the often much larger ongoing EEG or background activity. According to the most widely accepted model, ERPs are signals generated by neural populations that are time-locked to the stimulus; these signals would be summed to the ongoing EEG activity. According to another model, however, ERPs are assumed to result, at least partially, from a reorganisation of the ongoing activity (Sayers et al., 1974). More recently, a study demonstrated that ERPs could be generated by stimulus-induced phase resetting of ongoing EEG components (Makeig et al., 2004).

The ERP waveform can be quantitatively characterised across three main dimensions: amplitude, latency, and scalp distribution. In addition, an ERP signal may also be analysed with respect to the relative latencies between its subcomponents. The amplitude provides an index of the extent of neural activity (and how it responds functionally to experimental variables), the latency (i.e. the time point at which peak amplitude occurs) reveals the timing of this activation, and the scalp distribution provides the pattern of the voltage gradient of a component over the scalp at any time instant (Sanei and Chambers, 2007).

The ERP signals are either positive, represented by the letter P, such as P300, or negative, represented by the letter N, such as N100 and N400 (see figure 1.4 on the next page for an example). The digits indicate the time in terms of milliseconds after the stimuli (audio, visual, or somatosensory). The amplitude and latency of the components occurring within 100 ms after stimulus onset are labelled oxogenous, and are influenced by physical attributes of stimuli such as intensity, modality, and presentation rate. On the other hand, endogenous components such as P300 are nonobligatory responses to stimuli, and vary in amplitude, latency, and scalp distribution with strategies, expectancies, and other mental activities triggered by the event eliciting the ERP. These components are not influenced by the physical attributes of the stimuli (Sanei and Chambers, 2007).
The ERP parameters such as amplitude and latency are the indicators of the function of the brain neurochemical systems. ERPs are also related to the circumscribed cognitive process. For example, there are interesting correlations between late-evoked positivities and memory, N400 and semantic processes, or the latencies of ERPs and the timing of cognitive processes. Therefore, the ERP parameters can be used as indicators of cognitive processes and dysfunctions not accessible to behavioural testing.

Nevertheless, there are overlapping components within ERPs, which represent specific stages of information processing, which are difficult to distinguish (Frodl-Bauch et al., 1999; Dien et al., 2003). An example is the composite P300 wave, a positive ERP component, which occurs with a latency of about 300 ms after novel stimuli, or task-relevant stimuli, which requires an effortful response on the part of the individual under test (Frodl-Bauch et al., 1999; Dien et al., 2003; Kok et al., 2004; Friedman and Cycowicz, 2001).

In the P300 wave (figure 1.4), the elicited ERPs are comprised of two main components: the mismatch negativity (MMN) and the P300 complex. The MMN is the earliest ERP activity that indicates that the brain has detected a change in a background of brain homogeneous events, and it has been detected even when the stimuli are unattended or ignored. Because the MMN is evoked by stimuli that fall outside the
focus of attention, it is considered to be a relatively automatic, pre-attentive response to stimulus deviance (Friedman and Cycowicz, 2001). The P300 complex represents cognitive functions involved in orientation of attention, contextual updating, response modulation, and response resolution (Dien et al., 2003; Kok et al., 2004), and consists mainly of two overlapping subcomponents P3a and P3b (Frodl-Bauch et al., 1999; Friedman and Cycowicz, 2001; Comerchero and Polich, 1999). P3a reflects an automatic orientation of attention to novel or salient stimuli independent of task relevance. Profrontal, frontal, and anterior temporal brain regions play the main role in generating P3a, giving it a frontocentral distribution (Friedman and Cycowicz, 2001). In contrast, P3b has a greater centroparietal distribution due to its reliance on posterior temporal, parietal, and posterior cingulate cortex mechanisms (Frodl-Bauch et al., 1999; Dien et al., 2003). The P3b wave is thought to reflect a higher-order violation of subject’s expectations of a given rule, constructed over a longer time period than the MMN, and has been closely linked to working memory (Goldstein et al., 2002; Polich, 2007) and conscious access (Dehaene et al., 2006; Dehaene and Changeux, 2011).

According to the classic view, ERP analysis is based on two basic assumptions: (1) the electrical response evoked from the brain is invariably delayed relative to the stimulus and (2) the ongoing activity is a stationary noise, the samples of which may or may not be correlated. Thus, ERP detection becomes a question of improving signal-to-noise ratio. A simple method to detect ERPs is to average across many repetitions (trials) with the main objective, of course, to increase the signal-to-noise ratio so that the EEG background activity is attenuated (Niedermeyer and Silva, 2004). Nevertheless, quantifying ERPs can be difficulty for several reasons, including environmental artifacts and intra-subject variability. This problem of classification can be solved using multivariate statistical methods (Donchin, 1969) and multivariate pattern (MVP) analysis King et al. (2013a).

1.4 Electroencephalography in DOC patients

Regarding EEG, we must differentiate between the ERP-based or active, passive and resting state methods used for the diagnosis of disorders of consciousness. The multidimensional construct of consciousness can be assessed through a variety of stimulation and recording paradigms, each one addressing particular aspects of consciousness across the spectrum of pathologies with disorders of consciousness.

For example, in case of coma state, Brainstem Auditory Evoked Potentials (BAEPs), Middle Latency Auditory Evoked Potentials (MLAEPs), Somatosensory Evoked Potentials (SEPs) and Visual Evoked Potentials (VEP) are general indicators of prognosis. The presence of these potentials indicate preserved brainstem, sensory pathways
and primary cortex function (Garcia-Larrea et al., 1992; Litscher, 1995). Nevertheless, these potentials are not related to cognition or high order function, hence they only indicate a bad prognosis with the absence of responses. Contrarily, cognitive ERPs are used to assess and predict higher order functions as language, attention and working memory.

The N400 (negative deflection 400 ms after the stimuli) ERP is found in healthy subjects when they see or hear semantically anomalous sentences like “The coffee is too hot to fly” versus “The coffee is too hot to drink” (Schoenle and Witzke, 2004), hence used a test to probe linguistic capabilities. When applied to DOC patients, the presence of an N400 due to semantic processing indicates preserved cognitive function and has only been detected in MCS patients (Rohaut et al., 2015).

Neural responses to motor imagery and spatial navigation tasks have proven to detect consciousness (Goldfine et al., 2011; Cruse et al., 2011, 2012) by asking subjects to imagine movements or navigate spatially and analysing the oscillatory responses. However, the specificity and sensitivity of this marker is still under discussion (Goldfine et al., 2013).

Another classic ERP used in DOC is the MMN (mismatch negativity, see 1.3.2 on page 12). In comatose patients, these evoked potential elicited by novelty detection (oddball paradigms) has been reported as a highly specific (>90%) predictor of awaking (Kane et al., 1993; Fischer et al., 2004; Naccache et al., 2005). Highly associated with the novelty P300, and with several flavours ranging from auditory beeps (Naccache et al., 2005) to the subject's own name (Fischer et al., 2010; Schnakers and Laureys, 2009), this ERPs has not proved to be always specific in regard to diagnosis of DOC in post-comatose states as MCS or VS/UWS: they have been detected in both MCS and VS/UWS patients (Faugeras et al., 2011; Naccache et al., 2005; Fischer et al., 2010).

The main reason for the specificity failure of the novelty P300 ERPs is that this particular complex represents several cognitive functions and consists of two overlapping subcomponents (P3a, P3b, see 1.3.2 on page 12). In 2009, Bekinschtein et al. 2009 proposed a two-level hierarchical oddball paradigm designed to differentiate the MMN and P3b responses (for details, see 1.5.2 on page 17). This new paradigm named Local-Global presented an effect when subjects were able to maintain conscious attention, thus implying capacity for conscious access. When applied to DOC patients, it was only detected in EMCS and MCS patients, with an exception of two VS/UWS patients that later recovered (Faugeras et al., 2012). Nevertheless, the main caveat of this test is its low sensitivity: it was detected in 53.8% of the EMCS patients and only 14.3% of the MCS patients.

Interestingly, a recent work proposed a multidimensional cognitive evaluation of DOC patients using ERPs (Sergent et al., 2017). During a 1.5 hour session, patients were probed for own name recognition, temporal attention, spatial attention, detec-
tion of spatial incongruence, motor planning and modulation of these effects by global context, reflecting higher-level function. This proof-of-concept study revealed that the combination of several ERP markers increased diagnostic sensitivity, particularly in the detection of minimally conscious states with the presence of high-level effects.

So far, these ERPs are mainly elicited by auditory, somatosensory and in less frequency, visual stimuli. The main reason behind this unbalance is the difficulty to force patients to open their eyes and fixate in the case of visual stimuli. An alternative stimulation is the use of Transcranial Magnetic Stimulation (TMS).

The Integrated Information Theory of consciousness (Tononi et al., 2016) states that, phenomenologically, each conscious experience is both differentiated (composed of several phenomenal distinctions that exist within it) and integrated (the distinctions are bound together in various ways). These properties, from a neurophysiological point of view, rely on the ability of multiple, functionally specialised areas of the brain to interact rapidly to form an integrated whole (Dehaene and Changeux, 2011; Friston, 2002; Laureys, 2005b; Tononi and Koch, 2008). In Information Theory, integration and differentiation is also defined as complexity. Under this premises, the spatio-temporal complexity of an EEG can act as a proxy to the combination of integration and segregation. Casali et al. 2013 proposed a marker of brain complexity that quantifies the response to direct cortical stimulation using TMS and measured by EEG, named Perturbational Complexity Index (PCI). In a follow-up study on a cohort of 38 MCS and 43 VS/UWS patients, authors reported a sensitivity of 94.7% and a specificity of 80% (Casarotto et al., 2016).

In a recent review, Bai et al. 2017 report several common aspects among various studies on resting state (or spontaneous) EEG recordings in DOC patients: spectrum power differences in alpha, delta and theta bands between MCS and VS/UWS patients; ratios between higher and lower frequencies correlates with CRS-R scores; spectral entropy, markers of EEG complexity and functional connectivity differentiates patients groups.

One particular work, analysed in this thesis, is the one presented by Sitt et al. 2014 which combines EEG-extracted markers from diverse theoretical frameworks and uses machine learning to predict the diagnosis of individual patients. In particular, they analyse dozens of markers including ERPs from the Local-Global paradigm and markers of information, complexity, connectivity and spectral power, obtaining an AUC of 78% when used to diagnose a cohort of 68 MCS and 75 VS/UWS patients.
1.5 Methods used in this thesis

1.5.1 The Coma Recovery Scale (Revised)

The current gold standard for the behavioural diagnosis of DOC patients is the Coma Recovery Scale - Revised (CRS-R). This scale, initially described in Giacino et al. 1991 and later revised in Giacino et al. 2004 was developed to characterise and monitor patients, detecting subtle but potentially meaningful changes in neurobehavioural function, while ensuring proper interrater reliability.

The CRS-R consists of 26 hierarchically arranged items that comprise 6 subscales addressing auditory, visual, motor, oromotor, communication, and arousal processes. Scoring is based on the presence or absence of specific behavioural responses to sensory stimuli administered in a standardised manner. The lowest item on each subscale represents reflexive activity, whereas the highest items represent cognitively mediated behaviours. Depending on the different scores in the subscales, the CRS-R will diagnose the patient as VS/UWS, MCS or EMCS.

1.5.2 The Local-Global paradigm

As described previously in 1.4 on page 14, the MMN and P3b events are close in time and extremely difficult to differentiate in individual subjects. For that purpose, Bekinschtein et al. 2009 propose a new paradigm named Local-Global which adds a second level of novelties to a classic oddball paradigm. A first level of regularities is defined at a local (or within trial) level, while the second level is defined at a global (or across trial) level. While disruptions of the local level regularities elicits the MMN and P3a ERPs, disruptions of the global regularities elicits a P3b ERP.

Each trial of the paradigm is formed by 5 consecutive sounds lasting 50 ms, with a 150-millisecond gap between the sounds’ onsets and an intertrial interval ranging from 1,350 to 1,650 milliseconds. The fifth sound can be either equal to or different from the first four; this defines whether the trial is standard or deviant at the local level. The second level of regularities is defined across trials (or at a global level); frequent trials (80%) define the regularity, and rare ones (20%) violate this regularity.

Two types of stimulation blocks are played to the subjects; in the XX blocks, the frequent stimulus corresponds to five equal sounds (local standard and global standard [LSGS]). In contrast, the infrequent stimulus corresponds to four equal sounds followed by a fifth different sound (local deviant and global deviant [LDGD]). In the XY blocks, the frequent stimulus corresponds to four equal sounds and a fifth different sound (local deviant and global standard [LDGS]). The infrequent stimulus corresponds to 5 equal sounds (local standard and global deviant [LSGD]). For more details see figure 1.5.
Figure 1.5: Illustration of the Local-Global paradigm. Each trial of the auditory paradigm was composed by 5 consecutive sounds. Four equal sounds define a local regularity (music notes). The fifth sound could be equal or different, defining a local standard or deviant trial, respectively. At a second level, frequent trials (80%, green shaded area) defined a global regularity and rare trials (20%, red shaded area) violated this regularity.

The local effect is quantified by contrasting all local deviant (LD) trials (LDGS+LDGD) versus all local standard (LS) trials (LSGS+LSGD). The global effect is quantified by contrasting all global deviant (GD) trials (LSGD+LDGD) versus all global standard (GS) trials (LSGS+LDGS). All subjects were presented with these four conditions, twice for each block type. Each block started with 20–30 global standard trials to establish the global regularity before the occurrence of the first global deviant trial. An example of the neural response to the Local-Global paradigm is shown in 1.6 on the next page.

Subjects were instructed to count the GD trials and asked after each stimulation block (3.5 minutes). For patients, if they appeared asleep, they were stimulated with pressure as recommended in the arousal facilitation protocol in the CRS-R.

1.5.3 Machine Learning

According to Mitchell 1997, “the field of machine learning is concerned with the question on how to construct computer programs that automatically improve with experience”. Throughout this thesis, we will focus on a specific type of learning named supervised learning, that is, algorithms that could learn from examples provided by a supervisor. In this case, the algorithm is first presented with a set of samples and its corresponding labels (training set). Once the algorithm has learnt from the examples, one can feed it with an independent set of samples (test set) and ask the algorithm to classify the samples based on the knowledge obtained from the training set. The performance of the algorithm can be assessed by comparing the labels provided by the algorithm and the ones provided by the supervisor. An good algorithm is the one who provides accurate labels within some reasonable error.

A classification algorithm can be univariate or multivariate. In the first case, a single variable or feature is provided for each sample. Multivariate Pattern Analysis
1.5. METHODS USED IN THIS THESIS

Figure 1.6: Local and Global effects as measured with EEG in one of the directors of this thesis. Timeseries of the Local Effect around the Fz electrode (top left) present the MMN around 140 ms after the onset of the 5th sound only when it is different from the first four. The associated scalp map (top right) shows a frontal/central negativity and a posterior positivity. Timeseries of the Global Effect around the Pz electrode (bottom left) depict the P3b component starting around 350ms after the onset of the 5th sound. The associated scalp map (bottom right) presents a central/posterior positivity.
(MVPA) classifiers refers to classification algorithms that uses more than one feature for each sample. Depending on the algorithm and the number of features, some of them could be discarded to reduce dimensionality. This is called \textit{feature selection}.

\textbf{Cross Validation}

In order to test the real performance of a classification algorithm, one must provide two sets of representative data from the underlying real population distribution. For the train set, it should be big enough so the algorithm learns generalizable rules and not particular and specific to the available samples. For the test set, its size will have a direct impact on the variance of the estimated performance.

A common problem we face when evaluating models in neuroimaging and clinical applications is that the number of available samples is severely limited. For example, data from DOC patients in the Pitié-Salpêtrière hospital in Paris is acquired weekly: 52 samples a year in the best case scenario. To address this issue, a widely used validation technique is to partition the available data into independent sets and use them for training and testing the algorithm. In order to reduce variability, this action can be repeated with different partitions. This method is called \textit{cross-validation} and there are several partitioning schemes. We here focus on two particular methods: Stratified Shuffle Split (also called Monte Carlo cross-validation) and Stratified K-Fold.

The Stratified Shuffle Split method consists on randomly shuffling the data and splitting the data into two partitions for training and testing the algorithm. The sizes of the partition are determined by the user, but both of them respects the label distributions of the original set. This action is repeated several times and the performance is estimated across the repetitions. A disadvantage of this methods is that there is no guarantee that a sample will be used for the validation set, and that validation sets overlap.

In the Stratified K-fold cross validation, samples are split into $K$ different groups (folds) that respects the original label distributions. The algorithm is then trained on $K - 1$ folds and tested on the remaining one. This action is repeated $K$ times with a different testing fold. An advantage of this method is that it guarantees that each samples is used for validation exactly once.

\textbf{Area under the ROC curve (ROC-AUC)}

Univariate and multivariate discrimination performance was summarised with Area Under the Curve (AUC) calculated from the receiver operator characteristic (ROC). For a binary classification system, the ROC pits the detection probability, commonly referred to as sensitivity against the probability of false alarm (1 - sensitivity). These probabilities are empirically estimated by moving the decision cut-off along the sorted
values of a continuous variable, e.g. a score, and evaluating its relation to the true label. In the case of traditional model-free univariate analysis, the score is the EEG-marker itself, in the case of univariate or multivariate machine learning it is the predicted probability of a given sample to belong to the target class. The AUC can then be conveniently used to summarise the performance, where a score of 0.5 is uninformative and equals to random guessing whereas a score of 1 amounts to perfect classification and 0 to total confusion, indicating negative correlation between the score and the label.

1.5.4 Statistics
Statistical analysis encompassed correlations using Pearson product–moment correlation coefficient (r) and Spearman rank correlation coefficient (rho) with corresponding probability values. Pearson chi-square and Wilcoxon rank sum test were used to test for independence between the diagnosis and the demographic information of the patients. Bayesian ANOVA was performed to test the differences between groups using the BayesFactor R package (JZS Bayes factor with “medium” default prior setting \( r = 0.5 \); Rouder et al. 2012; Morey et al. 2014; R Development Core Team 2016. Bayes factor (BF) interpretation was done according to the Kass & Raftery scale (Kass and Raftery, 1995). Differences between groups were also tested using 2-sided paired samples signed tests. Different between conditions at single subjects and group level were tested using paired t-tests. Performance of MVPA models was assessed using the Area Under the Curve (AUC). MVPA models were tested using nonparametric Kruskal–Wallis test adjusted for multiple comparisons. We extended our visualisations into hypothesis tests by employing the percentile bootstrap (Efron and Tibshirani, 1994). Accordingly, we generated 2000 bootstrap samples by drawing with uniform probability and replacement n samples from the dataset. The test-statistic of interest was then evaluated on each bootstrap sample. Two-sided 95% confidence intervals were obtained by querying the 2.5 and 97.5 percentiles and the significance-level was then obtained by inversion of the confidence interval that excluded the value under H0.

1.6 This work

1.6.1 Purpose
The purpose of this thesis is to analyse the current EEG-based tools used to diagnose disorders of consciousness with two major objectives: (1) validate and standardise the usage of state-of-the-art tools based on machine learning methods and (2) analyse and propose novel tests in order to increase the sensitivity of the current tool and improve the diagnostic accuracy.
1.6.2 Description of chapters

Chapter 2: Automation and identification of robust EEG-extracted markers for the diagnosis of DOC. As a first step, we analyse the validity and clinical usage of a previous publication which combines EEG-extracted markers and a Support Vector Machine classifier to diagnose DOC patients. In order to develop an online clinical tool, we automate the process and analyse the application across different clinical setups.

Chapter 3: Towards cross-modal integration as a measure of consciousness. Previous works on diagnosis of DOC using auditory ERP analyses show high specificity but low sensitivity. Past research on multimodal stimulation suggest the existence of a variety of neural responses that can be measured using EEG. In this chapter, we analyse the possibility of using the neural responses to multimodal stimulation and cross-modal integration for the diagnosis of DOC.

Chapter 4: Brain-Body interactions as a diagnostic marker for DOC. So far, all the assisting technologies used to diagnose DOC patients are based on a strict neuro-centric approach. Nevertheless, past research on healthy subjects demonstrate interactions between the body and the brain, which are also affected by cognitive processes. In this chapter, we shift away from this neuro-centric approach and analyse the possibility of evaluating DOC patients via the embodied paradigm, according to which body-brain functions contribute to a holistic approach to conscious processing.

Chapter 5: Summary and final remarks. In this chapter, we summarise the findings and discuss future lines of research.
Resumen en español

Introducción
El término conciencia ha sido estudiado desde disciplinas como la filosofía, psicología y ciencias biomédicas. Sin embargo, no existe en la actualidad una definición universal para ese concepto tan ambiguo (Baars, 2015). El estado de conciencia, es a su vez una construcción multidimensional (Bayne et al., 2016). Esto quiere decir que puede ser expresado en distintos niveles que describen el nivel de vigilancia, las funciones cognitivas y el estado corporal. Una de las aplicaciones clínicas en las que se requiere evaluar los niveles de conciencia es en el dominio de los desordenes de conciencia (Disorders of Consciousness, DOC). Los pacientes DOC están despiertos, pero no presentan signos de entendimiento acerca de sí mismo o el ambiente en el que se encuentran. En ciertos casos presentan signos de de actividad voluntaria, pero sin embargo siguen incapaces de comunicarse.

El diagnóstico de pacientes DOC tiene implicancia en varias dimensiones. Por un lado, los familiares y las personas cercanas a los pacientes quieren saber que es lo que pasa con sus seres queridos. En algunos casos, junto con los médicos, se deben tomar decisiones acerca del cuidado, los tratamientos e incluso la discontinuidad del soporte vital que mantiene al paciente con vida (Jox et al., 2012). En otro nivel, la sociedad requiere administrar eficientemente los recursos disponibles en hospitales y clínicas (Beecher et al., 1968) y el debate se centra en como se considera la conciencia en estos estados (Racine et al., 2008). En la actualidad, el diagnóstico se ha centrado en determinar de forma confiable una serie de etiquetas clínicas entre las que se destacan el estado vegetativo o síndrome de vigilancia no responsiva (VS/UWS) y el estado de conciencia minimal (MCS). Uno de las principales diferencias es que los pacientes MCS tienen un pronóstico más favorable que los pacientes VS/UWS.

A pesar de la especificación de los criterios para diagnosticar a los pacientes como MCS (Giacino et al., 2002), un estudio encontró que el uso de evaluaciones no estandarizadas presentan una tasa de error de 41% (Schnakers et al., 2009). Una de las herramientas estandarizadas es la escala de recuperación de coma en su versión revisada (CRS-R, Giacino et al. 2004). En la actualidad, la evaluación comportamental sigue siendo la herramienta más confiable. Sin embargo, avances en las técnicas de imágenes cerebrales han permitido documentar, calificar y cuantificar información acerca del estado y el pronóstico de los pacientes. Las tecnologías que permiten asistir a los médicos incluyen imágenes acerca de la estructura cerebral como la resonancia magnética (MRI) y la tomografía computada (CT). Por otro lado, se usan tecnologías de imágenes funcionales como la resonancia magnética funcional (fMRI, Demertzi et al. 2015), tomografía por emisión de positrones (PET, Stender et al. 2014) y electroencefalografía (EEG).
En particular, la electroencefalografía es una técnica que mide la actividad eléctrica que transcurre en el cerebro humano. Es una tecnología de fácil acceso y económica. La adquisición de EEG consiste en ubicar una serie de electrodos conectados a un amplificador. Los electrodos se ubican en el cuero cabelludo y capturan las diferencias de voltajes que resultan de la actividad neuronal.

Actualmente, para el diagnóstico de DOC, el EEG se utiliza de diversas maneras: con estímulos o en estado de reposo. Cuando se utilizan estímulos, se miden las reacciones neuronales ante distintos eventos. Estas reacciones se denominan potenciales evocados por eventos (ERP) y llevan un nombre de acuerdo a las características. Las dos más comunes en DOC son la mismatch negativity (MMN) que ocurre entre aproximadamente 100 ms después del estímulo, y la onda P300, un potencial positivo 300 ms después del estímulo. Sin embargo, el potencial MMN se ha percibido en pacientes en estado VS/UWS o coma (Faugeras et al., 2011). Por otro lado, el potencial P300 no es detectado en pacientes VS/UWS pero si en algunos pacientes MCS (Faugeras et al., 2012). El problema reside en que incluso en sujetos sanos, este potencial está presente sólo si el sujeto esta atento a la tarea (Bekinschtein et al., 2009). En cuanto a EEG durante un estado de reposo, se pueden analizar oscilaciones, complejidad y conectividad entre otras métricas. Un estudio particular, analizado en esta tesis, es el presentado por Sitt et al. 2014, que combina métricas extraídas del EEG usando aprendizaje automático y predice el diagnóstico individual de los pacientes. En la publicación original, reportan un area bajo la curva ROC (AUC) de 78%, obtenida en un grupo de 68 pacientes MCS y 75 pacientes VS/UWS.

El objetivo de este trabajo es analizar las herramientas disponibles para el diagnóstico de DOC con dos objetivos principales: (1) validar y estandarizar el uso de aprendizaje automático y (2) desarrollar nuevas evaluaciones y marcadores para incrementar la sensibilidad y exactitud de las herramientas actuales.

Descripción de los capítulos

Capítulo 2: Automatización e identificación de marcadores robustos para el diagnóstico de DOC extraídos de EEG. En un primer paso, analizamos la validez y la aplicación clínica de un trabajo previo. El estudio combina marcadores extraídos de grabaciones de EEG en un clasificador SVM para diagnosticar pacientes DOC. Con el fin de desarrollar una herramienta disponible a travez de Internet, automatizamos el proceso y estudiamos la aplicación en distintos contextos clínicos.

Capítulo 3: Hacia una medida de conciencia basado en integración cros-modal. Trabajos previos que utilizan estímulos auditivos y EEG para el diagnóstico
de pacientes DOC muestran alta especificidad pero baja sensibilidad. Por otro lado, estudios centrados en la estimulación multimodal (multisensorial) sugieren la existencia de distintas respuestas neuronales que pueden ser medidas con EEG. En este capítulo, analizamos la posibilidad de utilizar las respuestas neuronales a estimulaciones multisensoriales y la integración crosmodal (entre estímulos de distinto origen sensorial) para el diagnóstico de DOC.

Capítulo 4: Interacciones Cerebro-Cuerpo como un marcador para el diagnóstico de DOC Hasta el momento, todas las técnicas utilizadas para el diagnóstico de pacientes DOC se basan en un enfoque estrictamente neuro-centrico. Sin embargo, estudios en sujetos sanos han demostrado la existencia de interacciones entre el cerebro y el cuerpo, que a su vez son afectadas por los procesos cognitivos. En este capítulo, nos alejamos de este enfoque neuro-centrico y analizamos la posibilidad de evaluar los pacientes DOC a través del paradigma incorporado, en el que las funciones cerebro-cuerpo contribuyen a un enfoque holístico sobre el procesamiento consciente.

Capítulo 5: Resumen y conclusiones finales. En este capítulo, resumimos los resultados obtenidos y discutimos posibles investigaciones futuras.
Chapter 2

Automation and identification of robust EEG-extracted markers for the diagnosis of DOC

Parts of the work described in this chapter has been accepted for publication as “Robust EEG-based cross-site and cross-protocol classification of states of consciousness”. Raimondo F.*, Engemann DA.*, King JR., Rohaut B., Louppe G., Faugeras F., Annen J., Cassol H., Gossieres O., Fernandez Slezak D., Laureys S., Naccache L., Dehaene S., Sitt JD. Brain, to be published in Volume 141, Issue 11, November 2018.

Additionally, this work was the basis for a case report published as “Probing consciousness in a sensory-disconnected paralysed patient”. Rohaut B., Raimondo F., Galanaud D., Valente M., Sitt JD., Naccache L. Brain Injury, Volume 31, Issue 8, Pages 1-6. 2017.

Contents

2.1 Background .................................................. 28
2.2 Objectives .................................................... 29
2.3 Automation .................................................... 29
  2.3.1 Reasons to rewrite the code and automate the process .... 29
  2.3.2 Software Implementation ................................. 30
  2.3.3 Preprocessing ............................................. 30
  2.3.4 Optimisations and Performance .......................... 31
2.4 Replication .................................................... 31
  2.4.1 Methods ................................................. 34
2.1 Background

Over the last decade, several electrophysiological signatures of consciousness have been proposed, varying from simple quantifications of ERPs or oscillations, to complex topological summaries of connectivity. This chapter is based on one of the key studies in EEG-based DOC diagnosis (Sitt et al., 2014). This work analysed dozens of EEG-markers obtained from more than 150 EEG recordings of the Local-Global paradigm. The authors demonstrated that by using a Support Vector Machine (SVM) classifier combining several EEG-markers, the diagnostic precision was higher than any of the individual markers. Furthermore, when the clinical (behavioural) diagnosed VS/UWS patients where classified by the SVM as MCS patients, the proportion of those who later showed signs of consciousness significantly increased (Sitt et al., 2014).

The main result, a gain in diagnostic precision by combining several EEG measures, has both theoretical and clinical implications. Theoretically, the results indicate that the markers addresses distinct and dissociable features of conscious states. Clinically, they depict the usefulness of combining EEG measures for the diagnosis. Particularly, the increase in the probability of recovery of those clinically VS patients classified as MCS using the EEG markers, indicates that this approach is discovering information
2.2 Objectives

In this chapter, we will focus on the follow-up analysis after the work published in Sitt et al. 2014. The purpose of this work has both clinical and scientific objectives:

1. Evaluate the use of the previously developed model for the diagnosis of DOC in a clinical environment.

2. Provide a semi-supervised algorithm to obtain the results, in which there is no need for human intervention.

3. Analyse the implications of modifying data acquisition conditions in order to assess the use of the developed tool across clinical centres with heterogeneous EEG systems and acquisition protocols.

4. Obtain insight from the classification model and determine which are the markers that drives the distinction between VS/UWS and MCS patients.

2.3 Automation

2.3.1 Reasons to rewrite the code and automate the process

The development of a clinical tool that goes beyond research requires a high grade of attention to the software development processes. For example, a defect in a news website or word processor application could result on the inability to perform a certain task, the loss of data or a security breach. Nevertheless, the magnitude of this events do not directly compare to a defect in a software that predicts the diagnosis of a patient. The clinicians might take end-of-life decisions upon the wrong results due to an error in the software. It is of uttermost importance to ensure a high quality software through the development process.

Particularly in data handling for machine learning applications, it is crucial that the features are well defined and obtained in an objective manner, with no prior knowledge of the output to be predicted. When it comes to EEG, one of the first steps is to identify non-neural signals or artifacts. In addition to the neuronal activity, EEG captures the electrical activity of muscles, cardiac activity, movements and environmental noise. A common procedure is to visually inspect the recording and discard the portions of data
that are contaminated by any of this artifacts. This procedure is prone to be subjective, hence a potential problem is a bias in the extracted features.

To address these potential problems, the proposed solution is to create a high quality software which extracts the features in a fully unsupervised manner.

2.3.2 Software Implementation

The software was written in Python and C, using open-source libraries. All the EEG-measures described in Sitt et al. (2014) were re-implemented taking advantage of the already optimised Numpy and Scipy libraries for fast algebra and scientific computing (Jones et al., 2001). For general data processing and visualisation, we used the open source MNE software package (Gramfort et al., 2013, 2014). Machine learning was performed using the scikit-learn library (Pedregosa et al., 2012). Bash scripts and GNU-parallel (Tange, 2011) were used to distribute processes and obtain results in a faster way.

2.3.3 Preprocessing

The first procedure after data acquisition, and before the computation of ERPs and EEG-extracted markers is called preprocessing. The objective of the preprocessing step is straightforward: convert raw and contaminated data into artifact-free data ready to analyse. Depending on the analysis to be done, the steps and tools used in preprocessing can vary.

Data was first filtered using a 0.5 Hz high-pass 6th order and a 45 Hz low-pass 8th order FFT-based Butterworth filters. The second step consisted on epoching (separate in trials) from -200ms to 1336ms relative the onset of the first sound of the Local-Global paradigm.

We then used an adaptive outlier detection algorithm specifically developed to detect and reject contaminated electrodes and epochs. This adaptive algorithm first selects bad electrodes where more than 50% of the epochs present a peak-to-peak amplitude higher than 100 $\mu$V. The second step consists on computing the variance of each individual channel and its corresponding z-score across all channels. Channels with a z-score greater than 4 are discarded. This operation is repeated 4 times. The remaining data is then analysed at the epoch level: epochs with more than 10% of the channels outside the 100 $\mu$V peak-to-peak amplitude range are then discarded. Finally, the second step is repeated, but with the standard deviation of the channels filtered with a 4th order Butterworth high-pass filter at 25 Hz.

In order to use the same set of electrodes for every patients, electrodes marked as “bad” by the outlier algorithm are interpolated using a spherical spline interpolation.
Data was finally re-referenced using an average reference and baseline corrected over the first 200 ms window preceding the onset of the first sound.

The development of the peak-to-peak amplitude rejection algorithm led to the development of an automated algorithm for rejection and repair of bad trials in EEG and MEG signals. This algorithm estimates the individual peak-to-peak threshold for each channel, rather than M/EEG system-dependent user set threshold (Jas et al., 2016). The presented method capitalises on cross-validation in conjunction with a robust evaluation metric to estimate the optimal peak-to-peak threshold, extended to a more sophisticated algorithm which estimates this threshold for each sensor yielding trial-wise bad sensors. Depending on the number of bad sensors, the trial is then repaired by interpolation or excluded from subsequent analysis.

2.3.4 Optimisations and Performance

Among the EEG-extracted measures used, three of them needed a reimplementation in Python: Permutation Entropy (PE; Bandt and Pompe 2002), Weighted Symbolic Mutual Information (wSMI; King et al. 2013b) and Kolmogorov-Chaitin complexity (K; Sitt et al. 2014).

Due to the complexity of the algorithms, optimisations were done in C using OpenMP for multithreading, with its corresponding Python bindings. This allowed to perform the computation of all the markers for a single subject in about 30 minutes in a 16 Gb RAM Intel Core i7 type workstation.

These benchmarks are particularly relevant for the practical purpose of the system. The current implementation facilitates the computation of reference models that are estimated on EEG-measures from hundreds of clinical recordings to predict unseen patients. This not only facilitates more frequent updates of these reference models, which may be required for research purposes. It also lowers the maintenance burdens, i.e., of detecting and fixing software bugs.

2.4 Replication

To validate, both the results from Sitt et al. 2014 and our Python implementation, we computed all the markers and the main analysis of the previous study, using the same original data. After preprocessing, we analysed a final cohort of 98 patients with 142 recordings (see Paris 1 in 2.2 on page 42).

At the univariate group level, we obtained the same results. Figure 2.2 on page 33 shows the topographical maps originally published in Sitt et al. 2014 (Figure 2.1 on the following page) and the results obtained with the Python implementation.
Figure 2.1: Scalp topographies of the most discriminatory measures. The fifth column indicates whether the VS and MCS patients were significantly different (Black: \( p = 0.01 \), light grey: \( p = 0.05 \), white: not significant, uncorrected). The sixth column shows the statistics of a regression analysis of the measure across the four states of consciousness (VS < MCS < CS (EMCS) < healthy controls (H)). Black: \( p = 0.01 \), light grey: \( p = 0.05 \), white: not significant, uncorrected). Extracted from Sitt et al. 2014.
Figure 2.2: Reproduction of the figure from Sitt et al. 2014 using our Python implementation.
2.4.1 Methods

For the multivariate analysis, we computed the cross-validation (CV) accuracy using the markers extracted with our implementation. We extracted 28 EEG-markers as described in detail in Sitt et al. 2014. These markers can be grouped into four conceptual families, i.e., information theory, connectivity, spectral, and evoked response markers (See Table 1). In the original publication, additional markers were used but were omitted due to previously reported unsatisfying performance or redundancy. For a full list of markers and abbreviations see Table 2.1 on the next page.

For each marker, we extracted four summary statistics. Aggregations over epochs were done using an 80% trimmed mean and the standard deviation to account for the average and its fluctuations over time. Aggregations over electrodes were computed using the mean and standard deviation to account for the average and topographical fluctuations. Some markers presented additional dimensions which were aggregated using a mean or median, according to the corresponding literature. For example, we would first compute either the mean or the standard deviation across epochs, and then the mean or the standard deviation across sensors. Throughout the chapter we refer to these marker subtypes as “mean,mean”, “std,mean”, “mean,std” and “std,std” and in figures, for brevity, “m,m”, “s,m”, “m,s”, “s,s”. A description on the dimensions and the procedure can be seen in Figure 2.3 on page 36.

These markers were then fitted into a classifier trained to distinguish between the VS/UWS and MCS patients. The classification algorithm used in the original publication consisted on a Support Vector Machine (SVM). The SVM-based classifier (SVC) aims at finding the optimal linear combination of features ($w$) that separates the training samples with distinct classes in the hyperspace of features. A penalisation parameter is used to find a solution which is likely to generalise to another dataset, and hence avoid over-fitting. Here, the penalisation parameter C, was chosen by nested cross-validation among the values $= [.001 .01 .1 .2 .5 1 2 10]$ using a grid-search method (Pedregosa et al., 2012). The SVC can provide a continuous probability by fitting the distribution of the samples with regard to $w$ (Platt, 1999).

Additionally, we used the Extremely Randomised Trees (ET) algorithm (Geurts et al., 2006) to address a potential decrease of performance when generalising to new data. Unlike Support Vector Machines, Extremely Randomised Trees are non-parametric and robust by design, and are not sensitive to the measurement scale of the input data. This algorithm can handle so-called wide datasets in which more variables than samples are available. Moreover, it belongs to the family of adaptive algorithms capable of scaling the complexity of the learned model to the amount of data available. The algorithm achieves its efficiency by generalising the non-linear decision tree approach. Single decision trees are non-parametric rule-based models that automatise variable selection.
### 2.4. REPLICATION

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Marker</th>
<th>Family</th>
</tr>
</thead>
<tbody>
<tr>
<td>PE Θ</td>
<td>Permutation Entropy</td>
<td>information theory</td>
</tr>
<tr>
<td>K</td>
<td>Kolmogorov Complexity</td>
<td>information theory</td>
</tr>
<tr>
<td>wSMI Θ</td>
<td>Symbolic Mutual Information (weighted)</td>
<td>connectivity</td>
</tr>
<tr>
<td>α</td>
<td>Alpha PSD</td>
<td>spectral</td>
</tr>
<tr>
<td>∥α∥</td>
<td>Normalized Alpha PSD</td>
<td>spectral</td>
</tr>
<tr>
<td>β</td>
<td>Beta PSD</td>
<td>spectral</td>
</tr>
<tr>
<td>∥β∥</td>
<td>Normalized Beta PSD</td>
<td>spectral</td>
</tr>
<tr>
<td>δ</td>
<td>Delta PSD</td>
<td>spectral</td>
</tr>
<tr>
<td>∥δ∥</td>
<td>Normalized Delta PSD</td>
<td>spectral</td>
</tr>
<tr>
<td>γ</td>
<td>Gamma PSD</td>
<td>spectral</td>
</tr>
<tr>
<td>∥γ∥</td>
<td>Normalized Gamma PSD</td>
<td>spectral</td>
</tr>
<tr>
<td>θ</td>
<td>Theta PSD</td>
<td>spectral</td>
</tr>
<tr>
<td>∥θ∥</td>
<td>Normalized Theta PSD</td>
<td>spectral</td>
</tr>
<tr>
<td>MSF</td>
<td>Median Power Frequency</td>
<td>spectral</td>
</tr>
<tr>
<td>SE90</td>
<td>Spectral Edge 90</td>
<td>spectral</td>
</tr>
<tr>
<td>SE95</td>
<td>Spectral Edge 95</td>
<td>spectral</td>
</tr>
<tr>
<td>SE</td>
<td>Spectral Entropy</td>
<td>spectral</td>
</tr>
<tr>
<td>CNV</td>
<td>Contingent Negative Variation</td>
<td>evoked</td>
</tr>
<tr>
<td>P1</td>
<td>P100 evoked potential</td>
<td>evoked</td>
</tr>
<tr>
<td>P3A</td>
<td>P3a evoked potential</td>
<td>evoked</td>
</tr>
<tr>
<td>P3B</td>
<td>P3b evoked potential</td>
<td>evoked</td>
</tr>
<tr>
<td>GD-GS</td>
<td>full contrast</td>
<td>evoked</td>
</tr>
<tr>
<td>LD-LS</td>
<td>full contrast</td>
<td>evoked</td>
</tr>
<tr>
<td>LSGD-LDGS</td>
<td>full contrast</td>
<td>evoked</td>
</tr>
<tr>
<td>LSGS-LDGD</td>
<td>full contrast</td>
<td>evoked</td>
</tr>
<tr>
<td>∆P3A</td>
<td>Contrasted P3A (LD vs LS)</td>
<td>evoked</td>
</tr>
<tr>
<td>∆P3B</td>
<td>Contrasted P3B (GD vs GS)</td>
<td>evoked</td>
</tr>
<tr>
<td>∆MMN</td>
<td>Contrasted MNN (LD vs LS)</td>
<td>evoked</td>
</tr>
</tbody>
</table>

*Table 2.1: Acronyms used for EEG-measures*
Figure 2.3: Diagram of the dimensions for each of the families of markers. To compute the scalar values, dimensions were first aggregated by epochs (1), then by sensors (2), and finally, if needed, by the remaining dimensions. (3)
and can be thought of as learning a “regression surface” from the data by recursive orthogonal partitioning (Efron and Hastie, 2016). In other words, decision trees map joint value ranges of the input variables to values of the outcome variable. However, decision trees poorly generalise to new data. The Extremely Randomised Trees retains all benefits of decision trees while mitigating their excessive variance and poor generalisation capability. This is achieved by averaging over many randomly constructed, hence uncorrelated, decision trees. To avoid overfitting, we used 2000 trees, limited the tree depth to a value of four and to maximise randomisation and minimise masking effects in feature importances (Louppe et al., 2013), we constrained the maximum number of features used for finding split points to one.

For cross-validation, a group Monte Carlo sampling scheme was used with a training set size of 80 percent, a testing set size of 20 percent and 50 iterations. The Monte Carlo procedure is known to minimise estimation variance and has been shown to yield low positive cross-validation bias (Varoquaux et al., 2017). The group variant consisted in exclusively assigning subjects to either the test or the train sets in order further avoid positive bias due to intra-subject sample correlations that are known to inflate the performance and constitute a violation to the assumptions of generalisation inference.

2.4.2 Results

Using our automated implementation, we obtained an AUC of 78.32% with the SVM classifier and an AUC of 77.35% with the ET algorithm, consistent with the 78% AUC reported in Sitt et al. 2014.

2.5 Validation

Despite the successful replication of the results obtained in Sitt et al. (2014), there was still a methodological risk: with repeated analysis on the same data, one risks to overfit the classifier and overestimate its diagnostic accuracy. For this purpose, the recommended approach is to test the accuracy on unobserved data. We then analysed 107 recordings from 92 patients registered after 2014 (see Paris 2 in 2.2 on page 42).

2.5.1 Methods

To assess the diagnostic accuracy on new data, we contrasted the performance of the classifiers against empirically estimated chance levels. These values were obtained from dummy classifiers which generates random predictions based on simple rules that do not depend on the actual predictors but could nevertheless lead to correct guess. Common dummy classifiers consist in 1) predicting the most frequent class, and 2) stratified
random guessing while respecting the class distributions. We then used the bootstrap method to analyse the AUC and differences of the means. Following the recommendations by Efron and Tibshirani 1993, we generated 2000 samples by drawing with uniform probability and with replacement.

2.5.2 Results

The resulting performance of the SVM showed a mean AUC of 74.27% while the ET classifier presented a mean AUC of 72.91%. Both performances were significantly different from the chance level AUC of 50.01% as estimated by a dummy classifier (figure 2.4 on the facing page).

Even though the classifiers performed similarly, one key aspect is still unanswered: is the combination of markers better than any marker alone? In other words, the use of a multivariate classifier is still not tested against univariate solutions. The direct comparison between the univariate model-free AUC and multivariate models is not entirely correct. In the first case, we do not build a predictive model but test how 'good' each marker separates the MCS and VS/UWS patients. Indeed, the in-sample estimation of the univariate AUC might give positively distorted estimates of true classification performance. To use a directly comparable measure, we trained univariate versions of SVM and ET classifiers. This allowed us to use the identical classification and prediction framework as for our multivariate analysis and obtain predicted probabilities of DOC diagnosis from single markers.

Interestingly, both univariate SVM and ET depicted alpha power (mean, mean) as the best univariate marker with an AUC of 71.40% and 67.53% respectively. Nevertheless, both of them were under their respective multivariate mean (figure 2.5 on page 40).

2.6 Clinical application

2.6.1 Report

Once the markers were recomputed and validated, the next step was to use this procedure in the clinical settings. Having a trained classifier with an automated method could be useful to predict the diagnosis of patients based on the EEG-extracted markers. MNE-Python provides a useful reporting tool: a self-contained HTML file with images and text that can be easily transferred by mail and reviewed even without internet connection. This report provides quantitative information on the preprocessing results, individual markers values, ERP timeseries and statistics, as well as results from
Figure 2.4: Our new implementation, tested on unobserved data, presents an AUC of 74.27% with an SVM classifier and 72.91% with an Extremely Randomised Trees algorithm (top). When compared to a dummy classifier to estimate the empirical chance level, the results of the classifiers are significantly different (bottom). The results where computed using 2000 bootstrap replicates.
Figure 2.5: Multivariate classifiers outperform univariate counterparts. Both SVM (blue) and ET (red) multivariate classifiers presented a superior AUC than the respective univariate versions on $\alpha(m,m)$ (dashed).

the multivariate (SVM) and univariate (Logistic Regression) classifications (Figure 2.6 on the facing page).

2.7 Further steps: towards a universal tool

So far, we have provided strong evidence supporting our classification model for the automated diagnosis of DOC patients. Nevertheless, the domain of application is still restricted: all the data comes from the same clinical centre, acquired with the same EEG system and under the same conditions. If the purpose is to build a tool that can be used to assess disorders of consciousness, it is important to assess the behaviour of such procedures under different conditions.

In clinical applications, we identified three main sources of variability related to EEG-based diagnosis of DOC. First, the behavioural assessment, despite the standardisation efforts and guidelines, are performed differently. Second, EEG systems configurations varies from clinical centres: the amount of electrodes and positions, the sampling rate and the recording quality changes depending on the brand and model of the acquisition systems. Finally, every clinical centre performs the EEG acquisitions under its own protocols: stimulation paradigms or resting state with varying parameters, including the recording length.

To assess the possibility of creating a universal tool for the EEG-based diagnosis of DOC, a generalisation analysis must be carried out to address the potential limitations. What is the optimal duration for individual EEG-recordings? Which stimuli and task
2.7. **FURTHER STEPS: TOWARDS A UNIVERSAL TOOL**

Figure 2.6: A self contained HTML-report presents all the results from the automated processing. This includes quantitative information regarding the preprocessing procedures, markers, statistics and predictions.
CHAPTER 2. EEG-BASED DOC DIAGNOSIS

should the patient be exposed to, if any? How many EEG sensors should be used, and where should they be located? Can the same machine learning algorithm operate on recordings acquired in different clinical centres?

To answer these questions, we probed the robustness and generality of EEG-signatures of consciousness under several simulated and real conditions. We first analysed the potential use of the tool across different EEG-systems using simulated electrodes montages and recording length from the EEG data recorded at the Pitié Salpêtrière hospital in Paris, France. Finally, we analysed a cross-clinical centre analysis and cross-protocol generalisation using resting state EEG data from the Liège University Hospital in Belgium. A description of the dataset can be found in table 2.2.

<table>
<thead>
<tr>
<th>Protocol</th>
<th>Auditory Local Global Task</th>
<th>Resting State</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dataset</td>
<td>Paris 1</td>
<td>Liège</td>
</tr>
<tr>
<td>n(EEG)</td>
<td>142</td>
<td>78</td>
</tr>
<tr>
<td>n(patients)</td>
<td>98</td>
<td>78</td>
</tr>
<tr>
<td>n(UWS)</td>
<td>75</td>
<td>22</td>
</tr>
<tr>
<td>n(MCS)</td>
<td>66</td>
<td>57</td>
</tr>
<tr>
<td>Sex ratio (male/female)</td>
<td>2.0</td>
<td>1.2</td>
</tr>
<tr>
<td>Age (M [SD]), in years</td>
<td>46.5 [17.8]</td>
<td>45.4 [17.7]</td>
</tr>
<tr>
<td>Delay (M [SD]), in days</td>
<td>125.9 [372.9]</td>
<td>299.5 [823.6]</td>
</tr>
<tr>
<td>Delay (MD), in days</td>
<td>30.0</td>
<td>529.0</td>
</tr>
<tr>
<td>Delay (SD)</td>
<td>372.8</td>
<td>1227.6</td>
</tr>
<tr>
<td>Delay (min to max), in days</td>
<td>6 to 2611</td>
<td>11 to 5380</td>
</tr>
<tr>
<td>Anoxia (%)</td>
<td>29.6</td>
<td>N/A</td>
</tr>
<tr>
<td>Stroke (%)</td>
<td>29.6</td>
<td>N/A</td>
</tr>
<tr>
<td>TBI (%)</td>
<td>23.5</td>
<td>48.1</td>
</tr>
<tr>
<td>Other (%)</td>
<td>18.4</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Table 2.2: Description of the three datasets used throughout the analysis in this chapter. Paris 1 refers to the data used in Sitt et al. 2014 and in section 2.4 on page 31, Paris 2 to the data used for the validation analysis in section 2.5 on page 37. Data provided by the Liège university hospital was used in section 2.7.2 on the next page.

2.7.1 Generalisation to difference EEG recording configurations

Methods To assess the potential use across incompatible EEG systems, we downsampled the data spatially and temporally after the preprocessing step. Spatially, we selected standard EEG configurations with 6 different number of electrodes (256, 128, 64, 32, 16, 8). Position were selected such that they approximated realistic EEG caps respecting the international 10-20 system. Temporally, we reduced the amount of epochs
2.7. FURTHER STEPS: TOWARDS A UNIVERSAL TOOL

Figure 2.7: Alternative EEG configurations with 6 different number of electrodes (256, 128, 64, 32, 16, 8). Position were selected such that they approximated realistic EEG caps respecting the international 10-20 system. Temporally, we reduced the amount of epochs to 6 different percentages (1, 5, 25, 50, 75, 100), respecting the original proportion of trials.

Results When we compared the univariate in-sample performance of the markers from the cross-validated multivariate AUC using the SVM and ET classifiers, we observed that many individual markers exhibited dramatic variability of performance across different combinations of sensor and epochs configurations. Nevertheless, the multivariate classifiers fluctuated between 70.94% and 78.33% for the SVM and 72.08% and 77.76 % for the ET (figure 2.8 on the next page).

Although the overall performance of the multivariate classifiers were in similar ranges, the ET classifier seems to be more stable than the SVM with alternative spatial configurations (figure 2.9 on the following page). Both classifiers tended to improve as more epochs and sensors were used. For the ET classifier, optimal performance was achieved with 128 electrodes. Nevertheless, reasonable performance could still be obtained with only 16 electrodes and a minimum of epochs. On the other hand, the SVM classifier peak accuracy stands at 256 electrodes and 100% epochs, but stability is severely affected by the amount of epochs and sensors.

2.7.2 Generalisation to Resting State

When training the classifiers on the all available data from Paris (Paris 1 and Paris 2) but ignoring the markers derived from evoked responses (table 2.1) the ET achieved an
Figure 2.8: Multivariate classifiers perform robustly on alternative EEG spatial configurations and recording lengths. Across all 36 experiments (6 alternative sensors locations by 6 temporal subsamplings) in-sample markers AUC (gray) are less performant than multivariate models.

Figure 2.9: Performance of the ET is more stable across spatial and temporal configurations compared to the SVM. Both classifiers present a trend to better accuracy with more epochs and electrodes. Nevertheless, SVM presents higher variability when the number of epochs is reduced.
2.7. FURTHER STEPS: TOWARDS A UNIVERSAL TOOL

AUC of 78.28% on the Liège resting state data and the SVM an AUC of 81.62% (figure 2.10 on the next page). Both classifiers presented significant differences compared to a dummy classifier.

We then compared the univariate and multivariate generalisation performances. Interestingly, the best univariate performance was achieved with alpha power with the \((\text{mean}, \text{mean})\) variation in the ET and the \((\text{mean}, \text{std})\) variation in the SVM, with AUCs of 74.85% and 75.73% respectively (figure 2.11 on page 47).

Finally, and coming back to the EEG configuration generalisation analysis, we tested the performance of the models across the temporal and spatial subsamplings. We trained and tested our models using each one of the 36 EEG configurations (see 2.7 on page 43) independently, yielding a total of 1296 experiments \((36 \times 36)\). When trained on the Paris 1 dataset and tested on Paris 2, both multivariate models outperformed the best univariate ones. Mean AUCs were as follows: SVM AUC 72.30%, ET AUC 73.20%, SVM \(|\alpha(s,m)|\) AUC 69.58% and ET \(\alpha(m,m)\) AUC 70.99%. When trained on Paris and tested on Liège, the same effect was achieved: SVM AUC 75.97%, ET AUC 76.59%, SVM \(\alpha(m,m)\) AUC 73.47% and ET \(\alpha(m,m)\) AUC 73.04%. Nevertheless, as depicted before, best performances are achieved with the SVM classifier: 80.55% and 82.95% vs 78.77% and 81.20% obtained with the ET classifier for the Paris 2 and Liège test sets respectively. For a visual representation, see 2.12 on page 47.

2.7.3 Discussion

These findings show that robust generalisation can be achieved despite changes in the spatiotemporal configurations of the EEG and the recording protocols. We compared systematically two different multivariate models that always outperformed its univariate counterparts. While certain EEG-signatures, i.e., alpha band power and its fluctuations turned out to be useful as stand-alone classifiers we found that the advantage of multivariate over univariate classification was most striking when systematic differences between the training and testing sets were present. We showed that by relying on a robust classification algorithm, meaningful generalisation could be achieved even if the performance of individual markers varied systematically between datasets.

Our results demonstrate that diagnosis of DOC patients can be robustly inferred from multivariate pattern classification using a wide array of EEG configurations. This was also the case with a minimum of sensors and epochs and even when EEG configurations differed on the training and testing data, e.g., when training on 10% of the epochs with 8 sensors and testing on all epochs with 256 sensors. We observed that many individual markers were highly variable. Nonetheless, our models fluctuated between AUC scores of 71% and 78% for the SVM and 72% and 77% for the ET.

It is important to note though, that the choice between the SVM and ET classifiers
Figure 2.10: Both our models generalise to resting state. The ET classifier present an AUC of 78.28% and the SVM classifier an AUC of 81.62%. When compared to a dummy classifier to estimate the empirical chance level, the results of the classifiers are significantly different (bottom). The results were computed using 2000 bootstrap replicates.
2.7. FURTHER STEPS: TOWARDS A UNIVERSAL TOOL

Figure 2.11: Multivariate classifiers outperform univariate counterparts also when generalising to resting state data. Both SVM (blue) and ET (red) multivariate classifiers presented a superior AUC than its univariate versions on $\alpha(m, s)$ (dashed blue) and $\alpha(m, m)$ (dashed red).

Figure 2.12: ET classifier presents less variability than SVM across spatiotemporal subsamplings. All the different spatial and temporal configurations were considered for training and testing (1296 combinations, 36 for each set). The corresponding best univariate models were also considered next to the multivariate ones. The distribution of AUC scores is indicated by the histograms for both validation (left) and generalisation (right) sets. Samples are depicted underneath the histograms, with black solid lines indicating the mean of the distributions. Both multivariate classifiers outperformed the corresponding best univariate ones. In both cases, the ET classifier presented less variance, although the highest accuracy was always obtained using the SVM model.
CHAPTER 2. EEG-BASED DOC DIAGNOSIS

should be done based on the domain of application. While SVM reported higher AUCs for most cases, the variability across conditions was always lower with the ET classifier. In that sense, the ET is preferred as the model to *Fit Them All*, since it can absorb the heterogeneity of the datasets. On the other hand, if the priority is to maximise the AUC, then our recommendation is to maintain the data variability as low as possible and use an SVM classifier, since in homogeneous data outperformed the ET in most of the cases.

2.8 Insights from Machine Learning

It is possible to analyse, post-hoc, which are the markers that the models are using to perform the classification. In 2.7 on page 40 we showed that the Extra Trees classifier was robust and could serve as a model to *Fit Them All* since it tolerates the variability on the individual markers due to heterogeneous EEG spatial and temporal configurations. Given that the ET classifier trained on one configuration generalises above chance for other configurations, one can argue that this classifier is capturing the common substrate of the EEG markers that allows the distinction between VS/UWS and MCS patients, disregarding the spatial and temporal resolutions of the recordings. Hence, analysing the groups of forests can depict if such common substrate is related to one or several particular markers, or it is the combination and interaction among all of them what allows the model to maintain its performance.

2.8.1 Methods

To obtain this insight, we extracted the *variable importance* metric from the Extra Trees classifiers. When entropy is used as impurity criterion, this multivariate metric can be shown to correspond to a weighted average of the mutual information between one variable and the outcome, conditionally over any possible configuration of any subset of the other variables (Louppe et al., 2013). Moreover, it has been shown that in fully randomised trees variable importance is only driven by relevant variables and not uninformative ones (Louppe et al., 2013). Also, the variable importance can deviate systematically from the univariate AUC whenever information is shared between markers or the model has identified non-linear interaction effects. To enforce this interpretation, we used entropy as impurity measure and only used one single marker for splitting, which maximised randomisation and made the tree-growing independent from the data (Geurts et al., 2006; Louppe et al., 2013).
2.8.2 Results

Inspecting all 36 DOC-Forest classifiers trained on the Paris 1 dataset using different EEG-configurations, we observed that markers contributing most strongly on average belonged to different conceptual families (Figure 2.13 on the following page left). Specifically, permutation entropy and long-range connectivity also in the theta band as well as alpha frequency band power were top ranked, both, in terms of univariate discrimination and variable importance. In contrast, evoked markers, on average, often assumed values below 0.89%, which is less than would be expected if all markers were equally influential. We observed a positive but non-linear relationship between average AUC and average variable importance ($\rho_{\text{Spearman}} = 0.82$, $p < 0.001$). It can be seen that highly performing markers were disproportionally more important than expected for a linear association.

When we compared the models variable importance against each marker’s out-of-sample performance for the validation and generalisation cases (Figure 2.13 on the next page middle and right), we also found positive non-linear correlations in Paris $1 \rightarrow 2$ ($\rho_{\text{Spearman}} = 0.48$, $p < 0.001$) and Paris $\rightarrow$ Liège ($\rho_{\text{Spearman}} = 0.521$, $p < 0.001$). The display reveals that several univariate models showed reasonable generalisation performance with AUC values beyond .70. Highly performing markers were disproportionally more important for the DOC-Forest than would have been expected assuming a linear relationship. Again, these findings suggest that the DOC-Forest achieves generalisation by preferentially enhancing the influence of reliable markers.

2.8.3 Discussion

Our findings suggested that protocol-general markers were, overall, more reliable. Strikingly, these markers belong to different conceptual families were all related to neuronal dynamics in the theta and alpha range (Figure 2.13 on the following page). The robustness of these markers may be explained by the fact that no excessive averaging is needed for their extraction and their characteristic EEG-topographies are simple and easy to capture with few sensors. However, the tight relationship between variable importance and conditional mutual information (Louppe, 2014) suggests that these top-performing markers carry independent information. Indeed, recent research has suggested a rather complex picture of functional and pathophysiological landscapes. The complexity of theta-band signals and their long-range interactions could reflect distinct memory processes underlying consciousness, such as access and maintenance (Axmacher et al., 2010). Similarly, alpha band power may reflect global arousal and demands for dynamic inhibition required for functional encapsulation of cortical networks (For an overview see Sadaghiani and Kleinschmidt 2016). Moreover, intact con-
Figure 2.13: We observed a positive but non-linear relationship between univariate AUC and variable importance in the ET classifier. Left plot depicts the correlation between the univariate AUC and variable importance of the ET classifier when averaged across the 36 experiments. Middle and Right plots depict the same correlation for the validation and generalisation cases. Colours indicate the family of the marker, while the shape depicts the variant. The three correlations were statistically tested using Spearman rank correlation coefficient.

Consciousness has been related to the peak frequency of alpha and theta band oscillations originating from distinct cerebral generators (Schiff, 2010; Williams et al., 2013). This is further complicated by the fact that these generators can be selectively disrupted for different aetiologies and can show a variety of regional effects during anaesthesia (Purdon et al., 2013). While future experimental research is desirable to disentangle these facets, our findings suggest that the presence of independent physiological sources of information may enhance generalisation as it is unlikely that all of them are jointly corrupted on new data samples.

Markers of Evoked Potentials are subject to extensive neuroscientific validation and intuitively support clinical reasoning. The P3 markers, for example, belong to the most studied indices of consciousness in the EEG literature and are commonly used in brain computer interfaces settings (Lulé et al., 2013). They have been related to processing novelty in bottom-up information, the global neuronal workspace, access consciousness, and context-updating (Dehaene et al., 2006; Donchin and Coles, 1988; Pins and Ffytche, 2003; Polich, 2007; Sergent et al., 2005). Considering such markers for multivariate analysis may, thus, improve interpretability. Additionally, evoked markers indexing auditory novelty have been shown to be rather specific than sensitive (King et al., 2013a). Like-wise, it could be the case that candidate signatures of conscious access, e.g., P3b, may be more relevant to distinguish MCS+ from MCS- patients (Naccache, 2017). Although being deemphasised by the ET classifier, evoked markers may still...
have contributed positively. Indeed, excluding all evoked markers from the Paris 1 to Paris 2 generalisation actually reduced the performance marginally (AUC = 71%). One could, therefore, argue that, evoked markers should be considered for multivariate analysis applied to DOC whenever available, alongside a few robust markers.

2.9 Summary

In this chapter, we analysed and compared two multivariate classifiers for the diagnosis of disorders of consciousness. We demonstrated that electrophysiological signatures of consciousness can be robustly exploited across contexts and protocols by relying on robust machine learning techniques. We validated on unseen data, analysed generalisation to resting state EEG and analysed the individual marker contribution to the multivariate model.

The significant generalisation from task to resting state EEG deserves separate consideration. It is well conceivable that EEG signatures related to the functional axis of consciousness (Sergent et al., 2017), are accessible during task and resting state EEG. Accordingly, changing states of consciousness should impact markers of global house-keeping functions such as alpha band power, global long-range connectivity or signal complexity, irrespective of the context. For instance, for a patient with locked-in syndrome we observed EEG-patterns similar to healthy persons during rest (Rohaut et al., 2017). And indeed, we observed significant generalisation from task to resting state EEG.

These findings imply that EEG signatures of consciousness can be reliably extracted from different contexts and combined into coherent predictive models, encouraging future efforts in large-scale data-driven clinical neuroscience.
Durante la última década, se han propuesto varios métodos para la detección de conciencia, desde simples cuantificaciones de potenciales evocados u oscilaciones, hasta mecanismos complejos de análisis de topologías de conectividad funcional. Este capítulo se basa en un estudio clave para el diagnóstico de DOC a través de EEG. En Sitt et al. 2014 se analizan decenas de marcadores extraídos de más de 150 EEG grabados bajo un paradigma auditivo llamado Local-Global (Ver 1.5.2). Los autores demostraron que utilizando un clasificador SVM que combine estos marcadores se puede tener una precisión de diagnóstico más alta que con cualquier marcador en forma individual. Adicionalmente, mostraron que cuando este método diagnostica un estado MCS que contradice al estado VS/UWS diagnosticado por una evaluación comportamental, la proporción de pacientes que mostraban mejoras en el futuro aumentaba significativamente. Estos resultados proveen una base para el desarrollo de una herramienta para el diagnóstico de pacientes DOC.

En este capítulo, nos enfocamos en continuar el trabajo de Sitt et al. 2014 con objetivos científicos y clínicos: (1) Evaluar el uso del modelo para el diagnóstico de DOC en ambientes clínicos, (2) proveer un algoritmo semi-supervisado para obtener los resultados sin intervención humana, (3) analizar la posibilidad de modificar las condiciones de adquisición de datos con el fin de utilizar la herramienta en distintos centros clínicos con protocolos de adquisición heterogéneos y (4) obtener información del modelo de clasificación para determinar cuales son los marcadores y las interacciones entre ellos que determinan la diferencia entre los pacientes VS/UWS y MCS.

El primer paso en el desarrollo de una herramienta semi-supervisada, donde el procesamiento se realiza sin intervención de un humano, requiere del desarrollo de algoritmos automáticos de procesamiento. Para ello, desarrollamos un conjunto de librerías en Python para el pre-procesamiento (filtro y remoción de segmentos de EEG no-neuronales) y la extracción de marcadores. Los resultados obtenidos fueron idénticos al análisis original, validando tanto los resultados originales de Sitt et al. 2014 (Figura 2.1) como nuestra implementación (Figure 2.2).

En segundo lugar, teniendo en cuenta la aplicación clínica en ambientes heterogéneos (distintos sistemas de EEG), proponemos dos modelos de clasificación: (1) un Support Vector Machine (SVM) y (2) un clasificador Extremely Randomised Trees (ET). Para ambos modelos, se extrajeron de los registros de EEG, 28 marcadores descriptos en Sitt et al. 2014 (ver tabla 2.1). Para cada marcador, se extrajeron 4 valores con distintas funciones estadísticas, resumiendo epochs y sensores usando combinaciones de desviación estándar y medias (ver figura 2.3). Utilizando estos modelos, se obtuvo un AUC de 78.32% con el clasificador SVM y 77.35% con el clasificador ET, consistentes con los resultados obtenidos en Sitt et al. 2014.
Sin embargo, cuando uno evalúa modelos utilizando los mismos datos, se corre el riesgo de caer en una condición de overfitting, en donde el modelo aprende a clasificar los datos utilizados de forma eficiente, pero cuando se evalúa en una nueva muestra de la población, el clasificador obtiene una performance significativamente baja. Para evaluar si nuestros modelos no se encuentran en esta condición, utilizamos una nueva muestra de datos de 107 registros obtenidos a partir del año 2014 (ver Paris 2 en table 2.2). Para verificar la performance en estos nuevos datos, contrastamos los modelos contra los valores de chance obtenidos en forma empírica. Estos valores se obtienen utilizando clasificadores que generan predicciones aleatorias utilizando reglas simples que no dependen de los predictores sino de la distribución de las clases de los datos de entrenamiento. Algunos ejemplos de clasificadores dummy consisten en (1) predecir la clase mas frecuente o (2) predecir aleatoriamente respetando las distribuciones de clases. Cuando evaluamos los modelos, obtuvimos una AUC media de 74.27% para el SVM y 72.91% para el ET. Ambos valores significativamente distintos del nivel aleatorio de 50.01% estimado por el clasificador dummy (ver 2.4).

En cuanto al desarrollo de una herramienta universal directamente aplicable en situaciones clínicas, es importante evaluar el modelo bajo las diferentes condiciones que se pueden encontrar en situaciones clínicas reales. En particular, no se puede pretender que todos los centros clínicos utilicen el mismo sistema de EEG, y que realicen las grabaciones bajo el mismo protocolo de adquisición. Para ello, definimos una serie de experimentos en los que emulamos diferentes configuraciones espaciales y temporales de EEG (ver 2.7) y evaluamos la capacidad de utilizar el modelo en grabaciones sin estímulo auditivo.

Cuando comparamos los modelos univariados con los modelos multivariados (SVM y ET) bajo diferentes configuraciones espaciotemporales, observamos que los marcadores individuales presentaron una gran variabilidad en la performance. Sin embargo, los clasificadores multivariados fluctuaron entre 70.94% y 78.33% para el SVM y 72.08% y 77.76% para el ET (ver 2.8). En el caso de las grabaciones sin estímulos auditivos, obtuvimos un AUC de 78.28% usando el clasificador ET y un AUC de 81.62% con el clasificador SVM. Ambos presentaros diferencias significativas con respecto al clasificador dummy.

Estos resultados implican que se puede realizar una clasificación robusta a pesar de los cambios espaciotemporales y del protocolo de registro. Mientras que algunos marcadores funcionan suficientemente bien de manera individual, los modelos multivariados mejoran y resisten ante estas variaciones. La diferencia entre los modelos ET y SVM radica en el dominio de aplicación. Mientras que el ET resulto menos afectado por la variabilidad individual de los marcadores, el SVM presenta una performance superior en condiciones homogéneas.

Adicionalmente, es posible analizar post-hoc cuales son los marcadores en los que
el modelo se basa para realizar la clasificación. Dado que el clasificador ET resultó eficiente en condiciones heterogéneas, uno puede argumentar que este modelo captura el substrato de los marcadores de EEG que permiten la diferenciación entre pacientes VS/UWS y MCS. Cuando analizamos la importancia de las variables del clasificador ET, una medida similar a la información mutua entre la variable y la predicción, podemos ver que la clasificación se basa en la combinación proveniente de marcadores de información, conectividad en la banda theta y el poder espectral en la banda alpha.
Chapter 3
Towards cross-modal integration as a measure of consciousness

Contents

3.1 Background .................................................. 56
3.2 Hypotheses .................................................. 58
3.3 Methods ...................................................... 59
   3.3.1 Cross-modal Local-Global ......................... 59
   3.3.2 Subjects ............................................... 61
   3.3.3 EEG acquisition ...................................... 62
   3.3.4 Analysis .............................................. 62
   3.3.5 Group-Level analysis ............................... 63
   3.3.6 Statistics ............................................ 63
3.4 Results ..................................................... 63
   3.4.1 ERP analysis ......................................... 63
   3.4.2 ERP characterisation through MVPA ............ 64
3.5 Discussion .................................................. 64

In the previous chapter, we analysed a solution to address one of the main problems of the Local-Global paradigm as a tool for the diagnosis of DOC: it is highly specific, but not sensitive. If the test is positive, the patient is clearly able to maintain conscious attention and thus in a higher state of consciousness. Nevertheless, conscious but inattentive healthy controls presented a negative outcome for the test (Bekinschtein et al., 2009).
The proposed solution using machine learning increases its sensitivity, nonetheless, it requires a dataset in which all the samples have already been labelled. Such cohort of patients required 6 years (from 2008 to 2014) of data acquisition. And even if one is able to acquire such data, there still no error-free and objective gold standard (Schnakers et al., 2009; Wannez et al., 2017), and one risks to reason in a circular manner: i.e. building a tool to label patients that works as good as the experts.

Furthermore, it is often the case that patients presents sensorimotor impairments. The Local-Global test relies only on auditory stimulation, which could yield negative results in case of auditory impairments. In 2017, we reported a case in which the test failed, but the machine learning approach described in chapter 2 on page 27 detected signs of consciousness (Rohaut et al., 2017) on a patient with no responses to auditory stimulation.

In this chapter, we propose to address these issues by analysing neural correlates of multisensory integration in DOC patients, with the objective to create a more sensitive test, independent of a database of previously diagnosed patients, and able to be used in sensory impaired patients.

3.1 Background

The way we experience our environment is by ongoing interactions between our brains and our senses (Ernst and Bülthoff, 2004). The brain is able to assemble impressions of the outside world and, in conjunction with intrinsic cognitive processes (memory, language, executive function), is able to “know” that these images and patterns are being viewed by and belong to the self (Fabrega, 2000). The senses provide the link between the body and the environment. The synergy or interaction among the senses, and the fusion of their information content, is described as multisensory integration (Stein and Stanford, 2008).

As regards external sources, evolution has provided us with a specialised set of sensory organs, each of which is linked to multiple specialised brain regions. There are obvious advantages associated with having multiple senses: each sense is of optimal usefulness in a different circumstance, and collectively they increase the likelihood of detecting and identifying events or objects of interest. However, the greater advantage comes with the ability to fuse the information content of different senses. In this case, the integrated product reveals more about the nature of the external event and does so in a faster and more accurate manner than would be predicted from the sum of its individual contributors (Stein and Stanford, 2008).

Neuronal mechanisms of multisensory integration in the cortex have been studied and are nowadays well characterised from single neuron perspective to whole brain
function (Stein and Stanford, 2008; Clavagnier et al., 2004; Lütkenhöner et al., 2002; Senkowski et al., 2008; Stekelenburg and Vroomen, 2007; Karns and Knight, 2009; Driver and Noesselt, 2008). On the relation between multisensory integration and consciousness, previous studies focused on attention and conscious content (Hartcher-O’Brien et al., 2017; Talsma and Woldorff, 2005), and bodily self-consciousness (Blanke, 2012). At any given moment, only a limited amount of information is consciously accessed and defines the current conscious content, which is reportable by the perceiving subject. At the same time, many other processing streams co-occur but remain non-conscious (Dehaene and Changeux, 2011), such as subliminal stimuli, motor reflex or sensory analysis (Kouider and Dehaene, 2007). By manipulating visual perception and creating a disruption with somatosensory input, a previous study recreated the neurological condition of out-of-body experiences, a disturbance of bodily self-consciousness (Lenggenhager et al., 2007). A recent work studied the effects of multisensory integration when stimuli were presented below the threshold of perception, concluding that multisensory inputs, even outside of awareness, are integrated and affect the phenomenological content of bodily self-consciousness (Salomon et al., 2017). Collectively, these studies show that multisensory integration can be studied either by manipulating the way sensory stimuli are combined or how intense these stimuli are. Importantly, they indicate that multisensory integration is realised beyond reportability. Interestingly, though, no much is known about how the conscious state affects multisensory integration. This knowledge is important if we were to tackle the necessary conditions for subjective experience (or self) to happen. Essentially, an unsolved issue is whether our ongoing sense of self, which presents itself so clearly in typical wakefulness but seems to demolish in sleep and anaesthesia, is preserved even when we are unable to report upon it (Windt and Metzinger, 2007). Importantly for clinics, could patients in vegetative/unresponsive conditions be considered as retaining basic subjective experiences?

Past research on ERPs depicts enhanced neural responses to multimodal stimulation when compared to unimodal stimuli (Giard and Peronnet, 2006; Talsma and Woldorff, 2005; Teder-Sälejärvi et al., 2002; Fort et al., 2002). In a recent publication, Chennu et al. 2013 uses a variant of the auditory Local-Global paradigm adding laterality mismatches. the effect of contralateral deviants depicted stronger MMN responses, reflecting the integrative processing of patterns across both auditory cortices to detect laterality shifts, and larger P300s waves, indexing the greater amount of cortical activation generated by the rare shifts in laterality of tones.

Taken together, these results suggest the existence of a variety of distinct neural responses that can be measured using EEG. While they have been linked to conscious content and attention, the relation between the neural components of multisensory integration and state of consciousness remains unexplored. It is hereby proposed to add
cross-modal stimuli as an extra layer to the Local-Global paradigm which is expected to promote multisensory integration, and therefore increase the sensitivity to detect consciousness in various states.

3.2 Hypotheses

For the purpose of this study, we propose an extension of the Local-Global paradigm to include three stimulation modalities (auditory, somatosensory and visual) that contains unimodal (same stimuli) and cross-modal (two modalities) trials.

The following are the working hypotheses:

1. Unimodal within-trial effects will differ between stimulation modalities, as a result of the different cortical sensory pathways.

2. Unimodal Global effects (associated with conscious processing) will be present and indistinctively of the stimulation modality, as a result of higher cognitive function not attributed to sensory pathways. All stimulation modalities will induce the same ERP.

3. Cross-modal and unimodal within-trial effects will differ, depicting the differential neural responses to cross-modal stimulation.

4. Within-trial effects will be detected in some DOC patients. Not all MCS patients will show the effect, with less proportion for VS/UWS patients, as a result of restricted cerebral integrity.

5. Global effects will be detected only in patients with preserved levels of awareness, implying the presence of multisensory integration induced by cross-modal stimulation.

6. Effects to cross-modal stimulation are expected to lay in an intermediate level. They will be present in fewer patients as compared to the within-trial effect and in more patients as compared to the global effect. Importantly, the presence of this response will not imply the presence of global effects (conscious attention).

At a first stage, the study will focus on the additional stimulation modalities, i.e. the local and global effects in the unimodal conditions (hypotheses 1 and 2).
3.3 Methods

3.3.1 Cross-modal Local-Global

In contrast to the original paradigm presented in Bekinschtein et al. 2009 in which the two different stimuli are presented as different tones, and in order to keep the conditions equal across stimulation modalities, this new version of the stimulation paradigm uses laterality as the rule, as introduced in Chennu et al. 2013. In other words, local standard trials presents all five stimuli either on the left or right sides, while local deviant trials presents the fifth stimulus on the opposite side to the first four. Side is chosen randomly with 50% chance each and timings between stimulus and trials were as in the original version. As described in 1.5.2 on page 17, two types of blocks are defined. In the XX blocks, the frequent trials (80% of trials) correspond to five ipsilateral stimuli (local standard and global standard [LSGS]). In contrast, the infrequent stimulus (20% of trials) corresponds to four ipsilateral stimuli followed by a fifth contralateral stimulus (local deviant and global deviant [LDGD]). In the XY blocks, the frequent and rare stimulus are reversed: four ipsilateral stimuli followed by a contralateral one defines the global regularity (local deviant and global standard [LDGS]) which is violated by five ipsilateral stimuli (local standard and global deviant [LSGD]).

In order to differentiate the effects of this paradigm from the previously described...
Figure 3.2: Illustration of the modification of the Local-Global paradigm that uses laterality as the difference between stimuli instead of modality-specific differences. Each trial is composed by 5 stimuli over a 600 ms interval as in the original paradigm (see 1.5 on page 18). The first four stimuli are delivered to the same side defining the local regularity. The fifth stimulus can be ipsilateral or contralateral, defining a local standard or deviant trial respectively. At the global (or across trial) level, the regularities are defined by frequent trials (80%, green shaded area) and violated by rare trials (20%, red shaded area). The choice of side for each trial is done pseudorandomly, keeping a balance of 50% for each side.
3.3. METHODS

Figure 3.3: Illustration of the types of trials used in the Cross-modal Local-Global paradigm. Three different stimulation modalities were possible. For each recording, two modalities were selected, defining 9 types of recordings. The matrix illustrates all the possible combinations, with the trials corresponding to unimodal recordings in the diagonal.

...in 1.5.2 on page 17, we will define two main effects: the Laterality Mismatch and the Rule Mismatch effects. The Laterality Mismatch effect is quantified by contrasting the trials with a fifth contralateral stimuli (all local deviant trials; LDGS+LDGD) versus the trials with all ipsilateral stimuli (all local standard; LSGS+LSGD). The Rule Mismatch effect is quantified by contrasting the rare trials (all global deviant; LSGD+LDGD) versus the rare ones (all global standard; LSGS+LDGS).

In each recording (20 minutes), subjects were presented with these four conditions, twice for each block type (XX, XY, XX, XY). Each block started with 24 global standard trials to establish the regularity. Subjects were instructed to count the GD trials and asked to report this number after each stimulation block (4.5 minutes).

Stimulations were presented as sounds, vibrations or visual stimuli. Sounds were presented using insert earphones either to the left or right ears. Vibrations were generated by two Eccentric Rotating Mass motors attached to the wrists (one in each wrist) and controlled by two Texas Instruments DRV2605 haptic driver for independent behaviour. Visual stimuli was delivered using two independent 8x8 LED matrix placed within virtual reality goggles that isolated the left and right visual fields. All devices were controlled by an Arduino Zero microcontroller that executed the stimulation as previously described.

For each recording, two stimulation modalities were selected: one for the first four stimuli of each trial, and the second for the last stimulus of each trial. Depending on the selection of the stimulation modalities, recordings were either unimodal (equal modalities) or crossmodal (different modalities). A total of 9 different types or recordings were possible (figure 3.3).

3.3.2 Subjects

A group of 44 right-handed healthy volunteers (35/9 female/male, mean ± STD age is 25.20 ± 4.1) participated in the study. Inclusion criteria was set to individuals aged 18-80 with normal binaural hearing, no tactile or visual impairment and no history of neurological or psychiatric disease. Participants were recruited via the RISC system from the Centre Nationale de la Recherche Scientifique (CNRS) in France. Subjects gave...
written and consent to participate in the study and received a remuneration of €40. Each participant was subject to three recordings which shared the same stimulation modality for the last stimulus (one row of figure 3.3 on the previous page) during a one hour session.

### 3.3.3 EEG acquisition

Data were acquired using 256-channel high density EEG net and a Net Amps 300 amplifier developed by Electrical Geodesics. Data were preprocessed in Python using MNE-Python (Gramfort et al., 2013, 2014) and a self-developed python library. EEG data were filtered between 0.5 and 45Hz and epoched 200ms before and 1356ms after trial onset. Artefacts were rejected by visual inspection of all channels. Channels which did not record activity were excluded from further analysis and interpolated at a later stage. Artefacts originating from eye blinks, muscle movements and electrical interference were visually identified and removed using independent component analysis (Lee et al., 1999). Following the interpolation of missing channels, data were re-referenced to the average and baseline-corrected relative to a 200ms interval before the presentation of the last stimulus in the five-stimulus trial sequence. We then removed channels from the face and neck, retaining a total of 224 channels for further analysis.

### 3.3.4 Analysis

Evoked Responses (ERP) were computed by averaging all trials for each condition described in 3.3.1 on page 59. Additionally, to characterise the temporal dynamics of the observed effects, Time Generalisation decoding was performed as described in King and Dehaene 2014.

Time Decoding consists of training a MVPA classifier to separate between the trials conditions using the values in each electrode as features and trials as samples. At each time point \( t \), a linear SVM estimator is trained and tested using Stratified 5-Fold cross validation and the performance is measured using the ROC-AUC metric. The Time Generalisation decoding consists on testing each one of the previously described estimators at all the different times points. That is, train an estimator on time \( t \) and test the ability to classify trials at a a different time point \( t' \), so as to estimate whether the scalp pattern is similar between \( t \) and \( t' \).

Similarity of patterns across stimulation modalities were tested using Time Generalisation with different training and testing conditions. That is, each estimator is trained to differentiate trials \( A \) from \( B \) when using a modality \( X \), and its discrimination accuracy is evaluated on the same type of trials but with the stimulation modality \( Y \). In this case, the results reveals the similarities between scalp patterns that are transferred
from modality $X$ to $Y$.

### 3.3.5 Group-Level analysis

In order to perform decoding at the group level, it is not enough to use the averaged epochs of each subject, as the amount of samples for the decoding will be equal to the number of subjects ($N \leq 15$). To overcome this issue, trials were averaged in groups (meta-epochs) of 10 epochs for each subject. Each group of epochs was randomly sampled 50 times, creating 50 different meta-epochs per subject. Decoding performance was estimated on each one of the meta-epochs sets.

### 3.3.6 Statistics

Evoked Responses (ERP) contrasts and Time Generalisation decodings were statistically tested using nonparametric permutation cluster test (Maris and Oostenveld, 2007). The statistical function was set to paired t-tests for the ERPs and independent samples t-test for the decoding. The clustering t-threshold was set equivalent to $p < 0.05$ for the given number of subjects or samples respectively.

### 3.4 Results

#### 3.4.1 ERP analysis

We first analysed the Laterality Mismatch effect, by contrasting the trials with ipsilateral stimuli against the ones with contralateral stimuli. For the auditory modality, the time series differ with a central negative deflection around 100 ms, and a centro-posterior positive peak with a maximum at 200 ms after the onset of the 5th stimuli (Figure 3.4 on page 65 top). Somatosensory contralateral stimuli, on the other hand, depicted an earlier frontal negative deflection (50ms to 150 ms after the onset of the 5th stimuli) and longer central positivity with a maximum around 300 ms after the onset (Figure 3.4 middle). For the visual modality, a negative deflection was detected from 50ms to 200ms after the onset of the 5th stimuli and a positive centro-posterior positive activation from 200ms to 400ms after the onset of the 5th stimuli (Figure 3.4 bottom). Nonparametric permutation cluster test indicated that 2 of the 3 contrasts were significantly different with $p = 0.0002$ for the auditory modality, $p = 0.06$ for the somatosensory modality and $p = 0.01$ for the visual modality.

Correspondingly, we performed the same analysis on the Rule Mismatch effect, by contrasting the frequent and rare trials. For the auditory stimulation, the contrast depicted a long-lasting posterior positivity starting 300ms after the onset of the 5th sound,
CHAPTER 3. CROSS-MODAL INTEGRATION

till the end of the trial (Figure 3.5 on page 66 top). The somatosensory modality presented the same long lasting positivity, with an additional early frontal positivity around 200-250ms, which presents the characteristics of time-locked eye blinks (Figure 3.5 on page 66 middle). Finally, the visual modality presented the posterior positivity starting 400 ms after the onset of the 5th sound, and an additional frontal positivity from 0 to 300ms, also consistent with the pattern of eye-blinks (Figure 3.5 on page 66 center). Nonparametric permutation cluster test indicated that the 3 contrasts were significantly different with \( p = 0.0002 \) for the auditory modality, \( p = 0.0012 \) for the somatosensory modality and \( p = 0.0006 \) for the visual modality.

3.4.2 ERP characterisation through MVPA

As a first step, we characterised the spatiotemporal patterns of the laterality and rule mismatches for each one of the stimulation modalities. Using Time Decoding, we found that Laterality Mismatches presented short patterns between 175ms to 250ms after the onset of the stimuli when stimulated with sounds, from 100ms to 375ms when stimulated with vibrations and from 200ms to 400ms when stimulated visually (Figure 3.6 on page 67) left). For the Rule Mismatches, the patterns appeared later and were more sustained in time. From 250ms to 600ms for auditory stimulation, 350 to 700 ms for somatosensory stimulation and 400ms to 700ms for visual stimulation.

To compare the spatiotemporal patterns, we applied the Time Generalisation decoding using different training and testing modalities. When we tested the Laterality Mismatches, we found a decrease in the decoding accuracy to almost the chance level (Figure 3.7 on page 68). That is, spatial patterns that separate with high AUC the ipsilateral and contralateral trials differ across modalities. In contrast, for the Rule Mismatches, the spatiotemporal patterns of the decoding accuracy are preserved, with some temporal shifts (Figure 3.8 on page 69).

3.5 Discussion

In this chapter, we aimed at analysing the viability of creating a multisensory Local-Global test with the objective of increasing its sensitivity while providing an alternative for auditory impaired patients. We proposed a novel stimulation paradigm in which stimuli is presented with sounds, vibrations and lights, while keeping the two-level oddball structure of the original Local-Global paradigm. We then analysed the neural responses to the effects for each oddball level as measured on healthy controls. We found that the effect of the Laterality Mismatch (short-term within trial mismatch) presented distinct spatiotemporal patterns across stimulation modalities. On the other hand, the
3.5. DISCUSSION

Figure 3.4: Topographies and ROI plots for the Laterality Mismatches comparing ipsilateral and contralateral stimuli for auditory (top), somatosensory (middle) and visual (bottom) modalities. Topographies depict the values of the contrast (subtraction) between conditions for the 224 scalp electrodes. Time series represents the average values around Cz for each condition. The $t = 0$ for the x-axis corresponds to the onset of the 5th stimuli.
Figure 3.5: Topographies and ROI plots for the Rule Mismatches comparing standard and deviant stimuli for auditory (top), somatosensory (middle) and visual (bottom) modalities. Topographies depict the values of the contrast (subtraction) between conditions for the 224 scalp electrodes. Time series represents the average values around Pz for each condition. The $t = 0$ for the x-axis corresponds to the onset of the 5th stimuli.
3.5. DISCUSSION

Figure 3.6: While Laterality Mismatch decoding patterns differ across stimulation modalities, Rule Mismatch decoding patterns do not present differences. Each one of the 6 images depicts the decoding AUC for each training and testing time. Decoding patterns of Laterality Mismatch (left) present a significant high AUC between 100ms and 400ms after the onset of the stimuli with some differences between auditory (top), somatosensory (middle) and visual (bottom) stimulation modalities. In contrast, Rule Mismatches (right) present later and longer decoding patterns. Significant clusters ($p < 0.05$) are depicted in color.
Figure 3.7: Spatiotemporal patterns of neural activity in response to Laterality Mismatches are not transferable across stimulation modalities. Each row of images depict the decoding AUC when training and testing using different pairs of modalities. Significant clusters ($p < 0.05$) are depicted in colour.
Figure 3.8: Spatiotemporal patterns of neural activity in response to Rule Mismatches are similar across stimulation modalities, with specific temporal shifts. Each row of images depict the decoding AUC when training and testing using different pairs of modalities. Significant clusters ($p < 0.05$) are depicted in colour.
CHAPTER 3. CROSS-MODAL INTEGRATION

Rule Mismatch effect (across-trial mismatch) were similar and shared spatiotemporal patterns across the three stimulation modalities.

Our results are consistent with previous reports on ERPs. Despite the fact that the original auditory Local-Global used tones as a varying factor instead of laterality, the obtained results with auditory stimulation are consistent with Bekinschtein et al. 2009, Chennu et al. 2013 (classic ERP analysis) and King et al. 2014 (Time Generalisation decoding). Regarding visual ERPs, we also found later responses to stimuli when compared to auditory responses, as reported in Giard and Peronnet 2006; Talsma and Woldorff 2005; Teder-Sälejärvi et al. 2002.

The Time Generalisation decoding across modalities depicted distinct spatiotemporal patterns for the Laterality Mismatch effect. This can be explained by the fact that the different responses are generated by the sensory-specific cortices, resulting in modality-specific components, as described in Giard and Peronnet 2006.

On the other hand, the Time Generalisation decoding across modalities did not show differences for the Rule Mismatch effect, besides the delay in the visual response. This results can be attributed to the fact that the obtained response, a P3b potential, reflects higher-order violation of the subjects expectation, which are not directly linked to the stimulation modality and the sensory pathways, but to the task being performed. This results are coherent with previous work on the P3b component, which associates the P3b to working memory (Goldstein et al., 2002; Polich, 2007) and conscious access (Dehaene et al., 2006).

Taken together, this preliminary results suggest that at a first step, the sensory-specific cortical areas are activated, while the task is then resolved in a common manner, independently of the modalities involved. This results settles the basis for the study of cross-modal interactions in relation to states of consciousness. Future work applied under different states of consciousness is needed to determine whether somatosensory and visual Laterality Mismatches responses are also nonconscious as the auditory Local Effect (Faugeras et al., 2012; Bekinschtein et al., 2009). Interestingly, this paradigm also contemplates using more than one modality in order to perform the task. This raises the questions on how and when is the information of these two modalities merged, and how this processing is affected by the different states of consciousness.

All in all, we believe that these results settles the bases for a deeper study with scientific and clinical implications. Scientifically, it paves the way for the study of the relation between multisensory integration and consciousness. Clinically, it is a start in the development of novel clinical tool which can help overcome the limitations of the current auditory Local-Global paradigm when applied to auditory impaired patients (Rohaut et al., 2017).
Resumen en español

En el capítulo anterior, analizamos una solución al principal problema del test Local-Global para el diagnóstico de pacientes DOC: es altamente específico, pero no sensitivo. Si el test es positivo, el paciente es claramente capaz de mantener atención consciente. Sin embargo, un sujeto consciente pero inatento puede resultar negativo en el test (Bekinschtein et al., 2009). La solución propuesta utilizando aprendizaje automático aumenta la sensibilidad, sin embargo require un set de datos en los que las muestras han sido diagnosticadas. En el caso de Sitt et al. 2014, esto requirió 6 años (desde 2008 a 2014). Además, en el caso de que uno pueda adquirir dicha cantidad de datos, no existe aun un diagnóstico 100% preciso y sin errores (Schnakers et al., 2009; Wannez et al., 2017), y uno corre el riesgo de razonar de manera circular: i.e. construir una herramienta para diagnosticar los pacientes que funciona tan bien (y tan mal) como los expertos. En este capítulo, proponemos una alternativa para mejorar la sensibilidad, analizando las respuestas neuronales a la integración multisensorial.

La forma en la que vivenciamos nuestro ambiente es a través de interacciones entre nuestros cerebros y nuestros sentidos. La sinergia o interacción entre nuestros sentidos y la fusión de su contenido es lo que se describe como integración multisensorial (Stein and Stanford, 2008). Más allá de la vasta capacidad sensorial del ser humano, la gran ventaja es la habilidad de combinar la información de los distintos sentidos. Colectivamente, el ser humano tiene la capacidad crear un producto integrado que revela más del evento externo, y de forma más rápida y eficiente, que lo que puede ser predecido de la suma de las contribuciones individuales (Stein and Stanford, 2008). La integración multisensorial ha sido estudiada desde la actividad a nivel neuronal hasta la función cerebral completa. Colectivamente, estos estudios indican que existe integración multisensorial mas allá de la percepción consciente. Estudios previos usando EEG muestran distintas respuestas neuronales ante estímulos provenientes de distintos sentidos. Sin embargo, la relación entre estado de conciencia e integración multisensorial no ha sido estudiada aun. Es nuestra propuesta agregar estímulos cross-modales al test Local-Global, forzando la integración multisensorial.

El test propuesto se realiza utilizando 3 tipos de estímulos: auditivos, visuales y somatosensoriales. Cada bloque del test utiliza uno o dos tipos de estímulos, dependiendo de si es un bloque unimodal o crossmodal respectivamente. Las hipótesis de trabajo son: (1) las respuestas neuronales correspondientes a los efectos locales serán distintas entre los bloques unimodales de distintos estímulos, (2) las respuestas a los efectos globales no serán distinguibles entre tipos de estímulos, (3) va a haber un nuevo efecto cuando se contrasten las respuestas a bloques unimodales con los bloques crossmodales. En esta tesis, sólo nos vamos a enfocar en las hipótesis 1 y 2, en condiciones normales (sujetos sanos).
Un grupo de 44 sujetos sanos participaron del estudio. Cada participante fue sujeto a 3 registros de EEG durante los que fueron estimulados usando la versión multimodal del test Local-Global. Se realizaron dos tipos de análisis sobre los datos: (1) un análisis clásico de potenciales evocados (promediando todos los trials y contrastando las condiciones, descripto en 3.3.1, y (2) un método para caracterizar las dinámicas temporales de los efectos obtenidos, llamado *Time Generalisation*, descripto en King and Dehaene 2014. Este último método consiste en entrenar clasificadores multivariados para separar trials de dos condiciones distintas. En este caso, utilizamos este método para caracterizar las diferencias y similitudes entre modalidades de estímulos distintas. Por ejemplo, entrenando el clasificador para diferenciar los trials correspondientes al efecto local cuando el estímulo es auditivo, y verificar la performance del clasificador para los mismos tipos de trials pero con estímulos visuales. Los patrones espaciotemporales compartidos serán aquellos en los que los clasificadores puedan distinguir sin importar el tipo de estímulo.

Los resultados del análisis clásico de ERPs mostraron la detección de irregularidades intra-trial (análogo al efecto local de la versión clásica del test). Sin embargo, los patrones topográficos y temporales no resultan similares (figura 3.4). El análisis *Time Generalisation* confirmó los resultados, obteniendo una performance mas baja cuando se evaluó la similitud de patrones entre distintas modalidades de estímulo (figura 3.7). Con respecto a las irregularidades inter-trial (efecto global), los patrones, tanto en el análisis de ERP clásico (figura 3.5), como en el *Time Generalisation* se mostraron idénticos (figura 3.8).

Nuestros resultados son consistentes con trabajos previos en ERPs. La diferencia en los patrones intra-trial puede ser explicado por el hecho de que las distintas respuestas son realizadas por estructuras corticales específicas a cada modalidad sensorial (Giard and Peronnet, 2006). Por otro lado, el análisis no mostró diferencias en los efectos inter-trial. Esto se atribuye a la respuesta obtenida, un potencial P3b, refleja una violación de alto orden de las expectativas del sujeto, que no está vinculada al tipo de estímulo y cortezas sensoriales específicas, sino a la tarea que se realiza. Estos resultados son coherentes con trabajos previos en la componente P3b, asociada a procesos cognitivos como la memoria de trabajo (Goldstein et al., 2002; Polich, 2007) y el acceso conscientes (Dehaene et al., 2006). Estos resultados sugieren que en un primer paso, las estructuras corticales especificas para cada sentido son activados, pero que luego la tarea se resuelve de la misma forma, independientemente de donde proviene el estímulo. Esto crea una base para el estudio de las interacciones crosmodales y la relación con los estados de conciencia.
Chapter 4

Brain-Body interactions as a diagnostic marker for DOC


Contents

4.1 Background ............................................................ 74
4.2 Hypotheses ............................................................. 75
4.3 Baseline cardiac activity .............................................. 75
  4.3.1 Methods ........................................................... 75
  4.3.2 Results ........................................................... 77
4.4 Cardiac markers of cognitive processing ......................... 83
  4.4.1 Methods ........................................................... 83
  4.4.2 Results ........................................................... 85
4.5 EEG and EKG Multivariate Pattern Analysis .................... 85
  4.5.1 Methods ........................................................... 85
  4.5.2 Results ........................................................... 87
4.6 Discussion ............................................................. 89
4.1 Background

A common aspect of the assisting technologies currently used to evaluate DOC patients is that all of them adapted a strict neuro-centric approach. However, classic (Critchley et al., 2001; Craig, 2002) and more recent studies (Critchley et al., 2001; Gray et al., 2007; Park et al., 2014; Seth et al., 2012; Park and Tallon-Baudry, 2014) in healthy subjects demonstrate that brain modulation of peripheral body functions are affected by concomitant cognitive processes. This evidence make the ‘brain-body’ interaction a relevant and promising space to evaluate consciousness level in patients. Cardiac activity is one of such peripheral body signals that has been linked to cognitive processes. For example, ‘bradycardia of attention’ refers to the effect of heartbeat frequency deceleration when the subject is engaged in an active cognitive task (such as target detection or auditory odd-ball counting) (Lacey and Lacey, 1978). Depending on the stimulation inter-trial interval, this cardiac cycle slowing is reversed when, or after, the target is detected (Van Der Molen et al., 1983; Jennings and Wood, 1977). More recently, it was also shown that by quantifying the neural events locked to heartbeats, one could predict whether a subject would report a fast flashing visual stimulus as perceived or not (Park et al., 2014). In addition, during complex cognitive processing, such as when playing chess, the heartrate dynamics, as measured before players made a move, could predict the likelihood of them eventually committing an error (Leone et al., 2012). Heartbeat-evoked cortical responses were further shown to differ in auditory interoceptive learning tasks (Canales-Johnson et al., 2015). Taken together, these studies suggest a bi-directional interaction between brain and heart that can be modulated by cognitive processes.

As regards patients with DOC, previous work has shown that cardiac autonomic markers, such as Heart Rate (HR) and Heart Rate Variability (HRV), were markers of autonomic system malfunction (dysautonomia) in traumatic brain injuries (Baguley et al., 1999). Specifically, patients with low scores on the Glasgow Coma Scale had altered autonomic functions including tachycardia and low HRV (Baguley et al., 2006). More recently, the same markers have been used to differentiate between VS/UWS and MCS when patients’ cardiac activity was recorded during sleep or after noxious stimulation (Leo et al., 2016). However, the link between these autonomic markers and conscious stimuli processing in DOC patients remains unknown.

In this chapter, we aimed at (1) evaluating if the heart-brain interactions could characterise the state-of-consciousness of DOC patients and (2) determining if the EKG-extracted information could complement the EEG evaluations of the patients. For the first objective, we quantified the modulation of cardiac cycle during the Local-Global paradigm (Bekinschtein et al., 2009) (see 1.5.2 on page 17). For the second objective, we contrasted the performance of multivariate patient classification of the state-of-
consciousness at the single patient level using either only the EEG markers or using the combination of EEG and cardiac cycle modulation markers.

4.2 Hypotheses

We hereby defined two working hypotheses:

1. Patients overall cardiac activity (HR and HRV) will be as described in previous works (Leo et al., 2016).

2. Markers of cardiac modulation by cognitive function, tested with the Local Global paradigm (see 1.5.2 on page 17), are expected to be present under the same conditions as the neural responses measured with the EEG. That is, the local effect should not discriminate between the patients group (Faugeras et al., 2011) and the global effect should be present only in patients with preserved levels of consciousness (Faugeras et al., 2012; King et al., 2013a).

Additionally, in order to test whether EKG has additional information to the EEG, we will test if the combination of EEG and EKG-extracted features modifies the accuracy of an automatic diagnosis by machine learning. If the accuracy is higher, this will mean that the EKG-extracted information is partially independent and complements the EEG-extracted information.

4.3 Baseline cardiac activity

4.3.1 Methods

Subjects and Patients  Patients admitted for consciousness evaluation at the Neurological Department of the Pitié-Salpêtrière Hospital, Paris between February 2008 and April 2015 were included. The neurological evaluation of the patients’ disorders of consciousness was performed by trained clinicians using the Coma Recovery Scale–Revised (CRS-R) as described in 1.5.1. Behavioural evaluations were performed systematically before each EEG recording.

In the present chapter, we aimed at characterising the cardiac cycle in relation to the state of consciousness as a post-hoc analysis. Since no EKG was available during the EEG evaluations, EKG time series were obtained using independent component analysis (ICA) on the EEG recordings for each patient. The current analysis only used the temporal location of the R wave peaks.
From the 259 patients originally assessed with EEG (130 VS/UWS, 129 MCS), 132 patients (51%; 60 VS/UWS, 72 MCS) were rejected due to the lack of a clear EEG recording or EKG reconstructed source that produced at least 40 samples for each stimulation block type. There were no differences between the included and excluded patients in terms of diagnostic state ($\chi^2[1, n = 259] = 2.07, p = 0.15$) and sex ($\chi^2[1, n = 259] = 0.21, p = 0.64$). Included patients were older than excluded patients ($48 \pm 18$ vs $44 \pm 17$ years; $W = 6701, p = 0.04$), and more patients suffered from anoxic as compared to traumatic injuries in the included group compared to the excluded group ($\chi^2[4, n = 259] = 12.84, p = 0.01$).

A final cohort of 127 (49%) patients remained: 70 VS/UWS (20 females, mean age = $45 \pm 19$ years, range 17–80, 12 traumatic, 21 assessed in a chronic setting [ie, >2 months postinsult]), and 57 MCS (17 females, mean age = $52 \pm 16$ years, range 21–79, 13 traumatic, 17 assessed in a chronic setting). Patient groups did not differ in terms of gender ($\chi^2[1, n = 127] = 6.2e^{-31}, p = 1$), etiologies ($\chi^2[4, n = 127] = 9.4, p = 0.051$), and chronicity ($\chi^2[1, n = 127] = 2e^{-30}, p = 1$). MCS patients were older than VS/UWS patients ($52 \pm 16$ vs $45 \pm 19$; $W = 2435, p = 0.03$). No patient had any history of cervical spinal cord injury or symptoms of autonomic dysfunction (eg, hemodynamic instability, abnormal HRV) at the time of EEG recording.

**EKG Extraction from EEG** In the absence of direct recordings of cardiac activity, EKG was extracted from the EEG using Independent Component Analysis. The independent components (IC) corresponding to the EKG were selected by visual inspection based on the spatial and temporal representation of the QRS complex. Raw EEG data was first filtered using an 8th order low-pass Butterworth filter at 45Hz and a 4th order high-pass filter at 0.5Hz (Figure 4.1 on page 78; top). Secondly, we computed three different ICA decompositions:

1. FastICA (Hyvärinen and Oja, 2000) parametrised to obtain the components that explain 99% of the variance and computed from raw filtered data.

2. INFOMAX (Bell and Sejnowski, 1995; Lee et al., 1999) parametrised to obtain 256 components from raw filtered data and

3. INFOMAX in combination to artifact channels rejection. Individual channels were removed when the temporal variance was more than 3 standard deviations away from the mean of the variance of the rest of the channels.

The independent component with the EKG information was selected based on the time series and the weights' topographies by visual inspection (Figure 4.1 on page 78; bottom). The selected time series had to clearly contain the R-peak corresponding to
the QRS complex. The R-peak had to be easily detected by using a simple threshold. The corresponding topography had to concentrate the mixing weights on the frontal right and posterior left electrodes. These electrodes are located in the right cheek, left maxillary junction and underneath the left mastoid, as depicted by previous studies on cardiac electrical fields (Dirlich et al., 1997).

We then picked the algorithm that presented the clearest decomposition, usually the one with the highest rank in descending order of explained variance. Finally, R-peaks onsets were obtained automatically by the algorithm described in Elgendi 2013. Subjects for which the EKG component was unclear were excluded from the analysis. Exclusion criteria was set to any of: EKG reconstructed signal with no clear R-peaks, detection failure by the automatic algorithm, or a topography of corresponding weights with a mix of peripheral and central electrodes.

Baseline cardiac activity  
The overall heart rate (HR) was computed by averaging the differences between consecutive R-peaks (RR Intervals) during the whole recording. Following the method described in Deboer et al. 1984, heart rate variability (HRV) spectral variables were obtained by computing the power spectrum decomposition on the point events time series from the detected R-peaks. Power Spectral Density was estimated in whole recording using Welch’s method with 32768 samples (131.072s) per segment and 28672 samples (114.688s) overlap using a Hanning window. HRV variables were extracted from the sum of the spectral power in three frequency bands: 1) very low frequencies (VLF, range 0–0.04Hz), 2) low frequencies (LF, range 0.04–0.15Hz) and 3) high frequencies (HF, range 0.15–0.4Hz).

R-peak locked EEG evoked responses  
EEG recordings were filtered as previously described, segmented from -200ms to 600ms relative to the onset of the R-peak and baseline corrected using the 200ms long window before R-peak. Bad channels and trials were rejected based on peak to peak amplitude exceeding 100µV. Bad channels were interpolated. The remaining trials were averaged. We performed a group analysis and obtained the mean evoked response for each group, and contrasted the VS/UWS mean evoked activity to the MCS one. Statistics on EEG responses were done using non-parametric cluster corrected permutation test (Maris and Oostenveld, 2007).

4.3.2  Results

EKG extraction method validation  
To test the homogeneity of the EKG-related ICA decompositions between groups, we computed the mean IC weights across subjects for the selected components. A sensor-wise Bayesian t-test showed evidence for no difference in the weights between the MCS and VS/UWS groups (Figure 4.2 on page 80;
Figure 4.1: (A) Filtered EEG time series from 7 EEG sensors from one VS/UWS patient. (B) Corresponding time series of 7 ICA components extracted from the previous EEG recording and the respective weights topographies. The independent component with cardiac information is shown in red. Dotted lines represents the automatically detected R-peak.
4.3. BASELINE CARDIAC ACTIVITY

We then averaged the cardiac cycle locked to the QRS complex at the group level and the contrasted the obtained time series between clinical groups (Figure 4.2; bottom). A single channel cluster permutation test found only one significant difference \( (p = 0.017) \) between 184 and 344 ms after the R-peak, consistent with the location of the T wave. No difference was found in the QRS complex.

Finally, we aimed at ensuring that the results obtained in terms of cardiac cycle induced by the processing of the auditory stimulation paradigm were not a side-effect of the EKG extraction methodology. In other words, we focused on testing that EEG-ICA extraction methodology was not injecting relevant EEG related activity to the EKG extracted signal. For this objective, we compared pure EKG to EEG extracted EKG. We performed simultaneous EEG-EKG recordings in an independent group of 24 healthy subjects and 32 patients (14 VS/UWS, 18 MCS). We applied the same EKG extraction method previously described and obtained 12 (50\%) healthy subjects and 12 (37.5\%; 3 VS/UWS; 9 MCS) patients with both direct EKG and indirect EEG-extracted EKG. We contrasted the two corresponding EKG time series in each trial, by subtracting the timing of the R-peaks in the direct EKG signal from EEG-extracted signal (Figure 4.3 on page 81). A repeated measures Bayesian ANOVA was computed using the REKG-REEG time differences as the study variable; trial types (LSGS, LDGD, LDGS and LSGD) and clinical state as factors. All the models including the trial type as a factor presented positive evidence in favour of no difference \( (BF_{01} \geq 4.27) \). Furthermore, the model that tested the interaction between clinical state and trial type presented even stronger evidence of no difference \( (BF_{01} \geq 15) \). Given that the only information used in this study was the timing of the R-peaks (automatically extracted and analysed within subjects), the here presented validation results strongly suggest that no effect was induced by the adopted EKG extraction methodology.

**Heart Rate and Heart Rate Variability** Overall Heart Rate was similar in patients across the two diagnostic groups \( (BF_{10} = 0.73; \) Figure 4.4 on page 82 A). When patients with overlapping behavioural CRS-R scores (CRS-R=6 or 7; 10 MCS; 20 VS/UWS) were excluded from the analysis to match the population in previous works (Leo et al., 2016) there was evidence for faster heart frequencies in the VS/UWS group \( (BF_{10} = 8.80) \). In the VS/UWS group, a positive correlation was identified between the HR and the CRS-R scores \( (\rho = 0.27, p = 0.02) \). No such correlation was found for the MCS patients. Similarly, HRV markers were comparable in both diagnostic groups (Figures 4.4 on page 82 B-D), HRV high frequencies \( BF_{10} = 0.62 \); HRV low frequencies \( BF_{10} = 0.36 \); HRV very low frequencies \( BF_{10} = 0.21 \). In the VS/UWS group a positive correlation was identified between the CRS-R and the HRV markers in high frequencies \( (\rho = 0.40, p = 0.0007) \) and in low frequencies \( (\rho = 0.27, p = 0.02) \). No such correlations were identified for the MCS group in either frequency.
(A) Group IC Topographies

(B) Group EKG

Figure 4.2: (A) Mean weights topographies for each clinical group (left). A sensor-wise Bayesian t-test shows evidence for no difference in the topographies between groups (right). (B) Mean and standard error of the mean for each clinical group QRS complex from the ICA-extracted EKG. A single channel cluster permutation test indicated significant differences ($p = 0.017$) only between 184 and 344ms after the R-peak, consistent with the location of the T wave.
Figure 4.3: EKG and ECG extraction methods comparison. We evaluated two independent groups of healthy controls (n=12) and patients (n=12) using simultaneous EEG and EKG recordings. For each subject, EKG was also extracted using the described ICA method. We then computed the differences between each R-peak onset detected in the direct EKG and the corresponding R-peak detected using ICA (left). Right panel shows the mean difference and 95% CI for each type of trial and subject as measured in samples (1 sample = 4 ms). Using Bayesian ANOVA, we found no evidence for a difference as an effect of the trial type ($BF_{01} \geq 4.27$) and strong evidence for no difference for the interaction between the type of trial and the clinical state ($BF_{01} \geq 15$)
Figure 4.4: Cardiac autonomic markers show no difference between clinical groups. Lower CRS-R
scores in VS/UWS patients correlates with a faster and less variable cardiac cycle as a manifestation
of the overall deterioration of the clinical condition. Each panel depicts the cardiac marker values
(y-axis, Heart rate (A), Heart rate variability in high frequencies (B), in low frequencies (C), and in
very low frequencies (D)) for each patient (dot), categorised by clinical group (top, 70 VS/UWS, 57
MCS, 12 Healthy) and by Coma Recovery Scale-Revised scores (CRS-R, bottom - only for patients).
The Spearman’s regression line between the scores and the EKG-related markers for the VS/UWS
patients, indicates a significant positive correlation between the CRS-R score and RR interval (A),
Heart Rate Variability in high frequencies (B), and in low frequencies (C). Boxplots with interquartile
range, median (black line) and mean (dashed line) represent the distribution of data in both clinical
groups.
4.4 CARDIAC MARKERS OF COGNITIVE PROCESSING

R-peak locked EEG evoked responses In terms of evoked responses to the cardiac activity as measured by EEG, a sharp peripheral bipolar topography was observed at the R-peak for both clinical groups (Figure 4.5 A and B). Between 0 and 250ms after the R-peak, both groups presented topographies following the pattern of the cardiac field artifact (CFA; Dirlich et al. (1997)). A cluster-level permutation test revealed a single significant cluster ($p = 0.034$; Figure 4.5 on the following page C) located between 144 and 540ms after the R-peak, with two spatial patterns, one similar to the CFA associated to the T-wave between 144 and 340ms, and second central spatial pattern after 340ms.

4.4 Cardiac markers of cognitive processing

4.4.1 Methods

To evaluate potential phase shifts in the cardiac cycle associated to the processing of different types of auditory stimuli, two intervals temporally locked to the onset of the fifth sound were defined (Figure 4.6 on page 85): (1) the PRE interval: the interval between the heartbeat (defined by the location of the R-peak) preceding the onset of the auditory stimulation, and (2) the POST interval: the interval between the stimulus onset and the following heartbeat. All time intervals were then labelled according to the contained auditory stimulation following the Local-Global paradigm (XX block: LSGS or LDGD; XY block: LDGS or LSGD; see 1.5.2 on page 17).

Finally, In order to avoid using peaks without a clearly defined temporal association to a given heartbeat (and not the previous or following one), we restricted the analysis to the trials in which both the PRE and POST intervals were between 20 and 600 milliseconds. A mean of $520 \pm 150$ trials per subject were included while $135 \pm 100$ trials were rejected ($20 \pm 13\%$). A repeated measures Bayesian ANOVA was computed for each interval using the ratio of rejected trials as the study variable and the trial label and clinical state as factors. All the models including the clinical state factor presented evidence for no difference (PRE $BF_{01} \geq 2.35$; POST $BF_{01} \geq 2.61$). When the models included the trial type factor, the test showed strong evidence for no difference (PRE $BF_{01} \geq 39.19$; POST $BF_{01} \geq 45.17$).

To test if conscious processing of auditory regularities affects the ongoing cardiac activity, we analysed the PRE and POST stimulus intervals for each group of subjects in relation with the type of trials. For the Local effect, each subject mean of the PRE and POST intervals corresponding to LD trials were subtracted from the mean of the LS ones. Similarly, for the Global effect, the mean of the PRE and POST intervals corresponding to GD trials were subtracted from the mean of GS ones.
Figure 4.5: R-peak locked EEG evoked responses shows differences between clinical groups. (A) Mean EEG topographies for each clinical group time locked to the R-peak at 0, 100, 250, 350, 450 and 500 ms. (B) Contrast and statistical comparison of the R-peak evoked potentials between clinical groups. (C) The left panel shows the only significant cluster of a permutation analysis ($p = 0.034$), the electrodes composing the cluster are shown with white circles (left). The right panel shows the time series of the corresponding cluster (mean and standard deviation across subjects). Two main modulations are observed, (1) in the time window corresponding to the T wave, a left-posterior positivity and a right-frontal negativity; and, (2) a central electrode spatial pattern positivity after the T-wave (>350ms), suggesting differences in the brain processing of the heart activity between VS/UWS and MCS patients.
4.4.2 Results

There was no evidence for difference in cardiac cycle modulation between groups due to the processing of the Local regularities in either the PRE ($BF_{10} = 0.19$) or the POST ($BF_{10} = 0.19$) intervals (Figure 4.7 on the next page). Within the groups, neither the VS/UWS nor the MCS patients presented significant differences between LS and LD trials (sign-test LD-LS trials, VS/UWS $p > 0.7$, MCS $p > 0.2$). In the case of the Global effect (GS vs GD trials), there was no evidence of modulation difference between groups due to the global auditory processing in the PRE interval ($BF_{10} = 0.21$). On the contrary, in the POST interval, there was a strong evidence for a difference between the MCS and VS/UWS groups ($BF_{10} = 43.07$). This result is explained by a shortening of the POST intervals in the GD trials compared to the GS trials in the MCS patients (sign-test GD-GS trials, $p = 0.007$) and no difference between GD and GS in the VS/UWS patients (sign-test GD-GS trials, $p = 0.55$). The small sample of healthy controls ($N=12$) included in this study presented a pattern of results similar to the MCS subjects (although not statistically significant).

4.5 EEG and EKG Multivariate Pattern Analysis

4.5.1 Methods

In order to analyse the relevance and independence of the markers to the diagnosis of DOC, we used Multivariate Pattern Analysis in combination with wrappers algorithms for feature selection (Kohavi and John, 1997). This method consists on training classifiers with different set of features and comparing the obtained performance. Based on the performance comparisons, a set of features can be defined as (1) strongly or weakly
Figure 4.7: Violations of global regularities induce cardiac cycle phase acceleration only in minimally conscious patients. Local violations did not affect the ongoing cardiac activity for the intervals between the stimulation onset and the preceding R-peak (PRE, top left) nor the following R-peak (POST, top right). Similarly, global violations did not affect the ongoing cardiac activity at the PRE interval (bottom left). In clear contrast, they induced shortened POST intervals (bottom right) only in the minimally conscious state (MCS) group (between-group contrast $BF_{10} = 43.07$; within-group sign-test $p = 0.007$). The small sample of healthy controls included in this study presented a pattern of results similar to the MCS subjects (although not statistically significant). Each dot represents a patient in vegetative state/unresponsible wakefulness syndrome (VS/UWS, $N=70$), in minimally consciousness state (MCS, $N=57$) or a healthy control (Healthy, $n=12$). Boxplots with interquartile range, median (black line) and mean (dashed line) represent the distribution of data in the clinical groups.
relevant when they are partially independent and contribute to an optimal classification or (2) irrelevant, when they do not contribute to the classification.

Multivariate pattern analyses were done using 120 EEG-extracted markers (corresponding to quantification of power spectrum and complexity in individual EEG sensors and information sharing between EEG sensors) as described in Sitt et al. 2014 (see chapter 2 on page 27) and 8 EKG extracted markers: HR, HRV (high, low and very low frequency), Local PRE, Local POST, Global PRE and Global POST. We trained a Support Vector Classifier (SVC) to distinguish between the VS/UWS and the MCS patients with a penalisation parameter (C) equal to 1. The SVC was repeatedly cross-validated with randomised stratified k-folding (k=8). Previously to the training of the classifier, relevant features were automatically selected keeping the highest 20% of the ANOVA F-value scores. Performance of the classifier was measured using AUC scores.

We defined 3 sets of features: (1) EKG markers of cognitive processes, corresponding to the PRE and POST intervals for the Local and Global contrasts (termed $EKG_{cog}$), (2) EKG markers of baseline vegetative function (termed $EKG_{veg}$) corresponding to the HR and HRV in the three frequencies previously defined and (3) EEG markers. We estimated the accuracy of the classification algorithm with 6 different combinations of these sets of markers: (1) EEG+$EKG_{cog}$+$EKG_{veg}$, (2) EEG+$EKG_{cog}$, (3) EEG+$EKG_{veg}$, (4) EEG markers only, (5) EEG with both $EKG_{cog}$ and $EKG_{veg}$ markers shuffled and (6) $EKG_{cog}$+$EKG_{veg}$ markers only. To minimise the effect of the random selection of folds, the AUC scores were averaged across 250 repetitions.

4.5.2 Results

Correlation between EEG and EKG markers of consciousness  We first tested the relationship between cardiac cycle modulation markers and EEG markers that previously were reported to distinguishing VS/UWS and MCS patients in Sitt et al. 2014. The modulation of the POST interval due to the Global Effect significantly correlated with EEG Kolmogorov Complexity ($K; r = -2.31, p = 0.02$), Permutation Entropy ($PE; r = -2.63, p = 0.01$), Spectral Entropy ($SE; r = -2.3, p = 0.02$), Weighted Symbolic Mutual Information ($wSMI; r = -0.19, p = 0.02$) and normalised Delta Power ($r = 0.2, p = 0.02$). No correlation was found between EEG evoked responses to the Global Effect and the phase shifts computed in the EKG (See 4.1 on the next page 1 for all markers). Nevertheless, none of the computed correlations survived a false discovery rate correction from multiple comparisons.

Multivariate patient classification by means of EKG and EEG markers  To determine if the EKG extracted information is partially independent to the consciousness related information extracted from the EEG, we trained classifiers to distinguish
<table>
<thead>
<tr>
<th>Marker</th>
<th>R Statistic</th>
<th>p value</th>
<th>p value (FDR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PE Θ</td>
<td>-2.63</td>
<td>0.009 *</td>
<td>0.138</td>
</tr>
<tr>
<td>K</td>
<td>-2.31</td>
<td>0.022 *</td>
<td>0.138</td>
</tr>
<tr>
<td>wSMI Θ</td>
<td>-2.34</td>
<td>0.020 *</td>
<td>0.138</td>
</tr>
<tr>
<td>α</td>
<td>-1.30</td>
<td>0.192</td>
<td>0.412</td>
</tr>
<tr>
<td></td>
<td></td>
<td>α</td>
<td></td>
</tr>
<tr>
<td>β</td>
<td>-1.91</td>
<td>0.058</td>
<td>0.251</td>
</tr>
<tr>
<td></td>
<td></td>
<td>β</td>
<td></td>
</tr>
<tr>
<td>δ</td>
<td>0.85</td>
<td>0.393</td>
<td>0.562</td>
</tr>
<tr>
<td></td>
<td></td>
<td>δ</td>
<td></td>
</tr>
<tr>
<td>γ</td>
<td>-1.67</td>
<td>0.096</td>
<td>0.321</td>
</tr>
<tr>
<td></td>
<td></td>
<td>γ</td>
<td></td>
</tr>
<tr>
<td>θ</td>
<td>-0.50</td>
<td>0.616</td>
<td>0.744</td>
</tr>
<tr>
<td></td>
<td></td>
<td>θ</td>
<td></td>
</tr>
<tr>
<td>MSF</td>
<td>-1.61</td>
<td>0.108</td>
<td>0.324</td>
</tr>
<tr>
<td>SE90</td>
<td>-1.87</td>
<td>0.062</td>
<td>0.251</td>
</tr>
<tr>
<td>SE95</td>
<td>-1.84</td>
<td>0.067</td>
<td>0.251</td>
</tr>
<tr>
<td>SE</td>
<td>-2.30</td>
<td>0.023 *</td>
<td>0.138</td>
</tr>
<tr>
<td>CNV</td>
<td>0.45</td>
<td>0.649</td>
<td>0.744</td>
</tr>
<tr>
<td>P1</td>
<td>-0.22</td>
<td>0.820</td>
<td>0.848</td>
</tr>
<tr>
<td>P3A</td>
<td>-0.86</td>
<td>0.387</td>
<td>0.562</td>
</tr>
<tr>
<td>P3B</td>
<td>-1.11</td>
<td>0.265</td>
<td>0.501</td>
</tr>
<tr>
<td>GD-GS</td>
<td>0.54</td>
<td>0.589</td>
<td>0.744</td>
</tr>
<tr>
<td>LD-LS</td>
<td>0.43</td>
<td>0.662</td>
<td>0.744</td>
</tr>
<tr>
<td>LSGD-LDGS</td>
<td>-0.31</td>
<td>0.756</td>
<td>0.810</td>
</tr>
<tr>
<td>LSGS-LDGD</td>
<td>-0.99</td>
<td>0.320</td>
<td>0.562</td>
</tr>
<tr>
<td>ΔP3A</td>
<td>-0.02</td>
<td>0.979</td>
<td>0.979</td>
</tr>
<tr>
<td>ΔP3B</td>
<td>-0.42</td>
<td>0.669</td>
<td>0.744</td>
</tr>
<tr>
<td>ΔMMN</td>
<td>-0.86</td>
<td>0.387</td>
<td>0.562</td>
</tr>
</tbody>
</table>

Table 4.1: Correlations between the Global Effect values as measured by the EKG POST interval and EEG markers used to diagnose the state of consciousness in DOC patients.
clinical groups and compared the performance of using as features EEG makers alone or combinations of EEG and EKG markers. Multivariate analysis combining EEG and EKG\textsubscript{cog} showed better performance compared to EEG and EKG\textsubscript{veg} markers and EEG markers alone (Figure 4.8 on the following page). Combining the EKG\textsubscript{cog} and EEG markers led to an improvement of the performance (EEG+EKG\textsubscript{cog}, AUC=76.1%; EEG+EKG\textsubscript{cog}+EKG\textsubscript{veg}, AUC=75.7%). On the other hand, when EKG\textsubscript{cog} markers were not included in the MVPA the performance did not differ from EEG alone (EEG only, AUC=73.7%; EEG+EKG\textsubscript{veg}, AUC=73.3%). As a control test for the effect of the number of features, classification was also computed combining EEG and label-shuffled EKG markers; in this case, the AUC was estimated at 73.6%. Using solely cardiac markers, the classifier performed above chance with a mean AUC of 60.1%. When we compared the performance of MVPAs that included EEG features, we only found significant differences when the MVPAs also included EKG\textsubscript{cog} versus when the MVPAs did not include these cardiac features ($p < 1e^{-9}$, Kruskall-Willis test, corrected for multiple comparisons). The inclusion of EKG\textsubscript{veg} features didn’t significantly changed the performance of the tested MVPA classifiers ($p > 0.1$).

**4.6 Discussion**

We here aimed at characterising consciousness state in patients with DOC by means of baseline heart activity and heart-brain interactions. We tested if cardiac-extracted information can complement single-patient EEG-based classification performance. When we contrasted behaviourally non-overlapping VS/UWS and MCS patients we found higher HR and HRV in the VS/UWS than MCS group, in accordance to Leo et al. (2016). This comparison included MCS patients who were in the higher end of the CRS-R scale versus the VS/UWS patients who were in the lower end of the CRS-R. When all DOC patients were included in order to retain clinical reality we did not find group differences of overall cardiac autonomic markers between the groups. This suggests a common underlying baseline cardiac function across patients. Interestingly, we found a positive correlation between CRS-R total scores and three autonomic markers (HR, HRV HF and HRF LF) only in the VS/UWS patients.

Our results are consistent with previous findings showing a relationship between the level of consciousness and dysautonomia in DOC after traumatic brain injuries. Specifically, low CRS-R scores were related to tachycardia in patients with low scores on the Glasgow Coma Scale (Baguley et al., 1999) and to lower HRV (in both high and low frequencies), which was considered as a symptom of a neurological disconnection syndrome (Baguley et al., 2006). Taken together, these results suggest that the diversity of behaviours characterising conscious states (associated with cortical processing) does
Figure 4.8: Cognitive EKG markers carries partially independent information from EEG. Six distinct multivariate classifiers were trained to distinguish between VS/UWS and MCS patients using different combinations of EKG and EEG markers. We used as features combinations of 120 EEG markers, cognitive EKG markers (EKG\text{cog}; PRE and POST intervals, Local and Global effects contrasts) and the vegetative function markers (EKG\text{veg}; Heart Rate and Variability). All the models that summed EEG markers and EKG\text{cog}, presented a significant increase in the classification accuracy (compared to MVPA of EEG without EKG\text{cog}, \( p < 1e^{-9} \)). Using only EEG markers (mean AUC 73.7\%) showed no significant difference with EEG in combination with EKG\text{veg} markers (mean AUC 73.3\%). As a control to equalise the number of features, the combination of EEG and all of the EKG markers with shuffled labels reported a mean AUC of 73.3\%. When we used only EKG markers, the classifier performed above chance, obtaining a mean AUC of 60.1\%. Means were estimated using 250 repetitions of stratified 8-fold cross validation. Each dot represents the mean value across folds for each repetition. Boxplots with interquartile range, median and mean (dotted line) represent the distribution of values for each set of features.
not necessarily translate into strong correlations with autonomic markers, such as HR and HRV. Therefore, the observed differences in these markers in VS/UWS patients on the lower end of the CRS-R scale seems to be associated with an overall deterioration of clinical condition, rather than to cognitive processing.

Our analysis of the heart evoked potentials revealed two results. First, we observed a statistical difference between VS/UWS and MCS in the CFA corresponding to the T wave but no difference in association to the QRS wave. The differences observed in the T-wave between VS/UWS and MCS patients, in the shape of a dipole with a left-posterior positivity and a right-frontal negativity, are similar to the reported cardiac repolarisation changes induced by mental stress (Gray et al., 2007) and neurodegeneration or stroke (García-Cordero et al., 2016). Although previous works depict a main modulation during the time window corresponding to the T-wave with frontal negativities, in our study the differences between the groups of DOC patients are highlighted by the cluster statistic in the posterior positive side of the dipole. Second, we found differences between VS/UWS and MCS patients in a time window after the T-wave. Crucially, this difference had a different topography to the previously described CFA. The maximal differences in the EEG were obtained in the central electrodes. Taken together these results further suggest differences in heart-brain interaction between VS/UWS and MCS patients.

In terms of cognitive processing, we analysed the cardiac activity while patients were evaluated with the Local Global paradigm aiming to probe cognitive-related responses on cardiac markers. Such brain-heart interactions have been previously shown in protocols where, by quantifying neural events locked to heartbeats, one could predict whether a subject would report a fast flashing visual stimulus as perceived or not (Park et al., 2014). In addition, during complex cognitive processing, such as when playing chess, the heartrate dynamics, as measured before players made a move, could predict the likelihood of them eventually committing an error (Leone et al., 2012). Heartbeat-evoked cortical responses were further shown to differ in auditory interoceptive learning tasks (Canales-Johnson et al., 2015) and emotional states (Couto et al., 2015). Taken together, these studies suggest a bi-directional interaction between brain and heart that can be modulated by cognitive processes.

In our protocol, we found that the cardiac cycle was modulated by the processing of global auditory regularities only in the MCS group. Specifically, MCS patients showed an acceleration of the timing of the heartbeat following the auditory stimulus (shortening of the POST interval) which disrupted the global regularity. No such modulation of cardiac cycle was found in the VS/UWS patients, nor any effect was found in either group for the local irregularities. No modulations of the PRE intervals were found, this suggests that the only observed modulation is a direct effect of the cognitive process of the stimulation. It is important to compare this results with previous works that
analysed the evoked responses in the EEG using the same protocol. These studies show that the violations of local regularities (in the form of a mismatch negativity response) can be detected in healthy and awake controls but also unconscious conditions such as subjects during sleep, coma and VS/UWS (Faugeras et al., 2011; Strauss et al., 2015; Morlet and Fischer, 2013). In contrast, disruptions of the global regularities (eliciting a P3b response) are only present in conscious and attentive subjects (although see Tzovara et al. (2015) and Naccache et al. (2015) for ongoing discussions). The fact that cardiac cycle modulation effect was present only associated to global irregularities (which requires maintaining conscious attention) and only in the MCS patients (who are generally characterized by more complex brain function compared to VS/UWS patients; Giacino et al. 2014) suggests that the source of this effect is a brain-driven indirect modulation due to the conscious processing of information.

A recent study demonstrated a link between conscious perception and cardiac activity in normal subjects (Park et al., 2014). Specifically, in visual detection task, subjects’ heart rate decreased during a warning cue and increased immediately after reporting the perception or not of the stimuli following the cue. When subjects responded correctly, following RR intervals were significantly shorter than the ones corresponding to an incorrect response. This indicates an interaction between conscious perception and the modulation of cardiac activity. Interestingly, previous studies showed that the characterisation of the modulation depends on the stimulation inter-trial interval. With short intervals, this cardiac slowing is reversed within the same cycle that the target is detected (Van Der Molen et al., 1983; Jennings and Wood, 1977). In our work we depict a shortening of the RR interval containing the stimuli. Nevertheless, only when the stimulus is known to produce neural modulations and only in patients with higher level of consciousness. Our attention-driven effect is consistent with these previous results and characterises the modulation in relation to the subjects’ overall level of consciousness.

Having a proficient test at the single-subject level is a clinical necessity in order to reduce the diagnostic uncertainty each case. The modulation of the heart cycle within each subject was not powerful enough to have a significant effect to distinguish the clinical state of individual subject. With the aim of improving the single case performance of diagnostics tests, and particularly in terms of EEG, we have shown that multivariate classification performance of the combination of 120 EEG markers (such as quantifications comprising connectivity analysis, information complexity, spectral analysis and evoked related potentials) outperformed the univariate classification accuracy, when markers were considered individually. This combination of EEG markers allowed an enhanced classification of conscious state at single-patient level (Sitt et al., 2014). Although the cardiac measures alone did not allow a single subject diagnosis, combining information from both neural and cardiac sources increased significantly the accuracy of
the classification of these patients. This indicates that the information extracted from
the modulations of cardiac activity due to cognitive processing is partially independent
from the neural correlates of consciousness as measured by EEG. To our knowledge,
this is the first time that body-related signals are considered as contributing factors in
data-driven diagnosis in patient with DOC. We think that such an embodied approach
to cognition (Clark, 1999) paves the way for further investigations of body-brain in-
teractions in DOC which might be informative not only for clinics but also for tracing
the neural correlates of consciousness. In the future, and with the aim of improving
the single case performance of this test, we will introduce novel versions of stimulation
paradigm (with stimulations contextually locked to the ongoing cardiac cycle).

In conclusion, we show a relation between autonomic nervous system function and a
stimulation paradigm exposing subjects to violations of auditory regularities in MCS pa-
tients. Our results suggest that cardiac cycle modulation is relevant for the assessment
of patients with DOC because it potentially carries partially independent information
when taken together with neural correlates of consciousness. We think that our work
opens a window to the study of DOC via the embodied paradigm, according to which
body-brain functions contribute to a holistic approach to conscious processing.
Resumen en español

Hasta este punto, todas las herramientas usadas para evaluar pacientes DOC se basan en un enfoque neurocéntrico. Sin embargo, estudios previos en sujetos sanos muestran que las modulaciones de las funciones periféricas, efectuadas por el cerebro, son afectadas por procesos cognitivos. La actividad cardíaca es una de las funciones periféricas que ha sido vinculada a los procesos cognitivos. Esta modulación, se ha detectado en forma bidireccional: e.g. la frecuencia cardíaca se ve afectada durante momentos de atención a eventos externos (Lacey and Lacey, 1978), y la percepción de estímulos visuales es afectada por el ciclo cardíaco (Park et al., 2014).

En este capítulo, nos enfocamos en (1) evaluar si las interacciones entre el cerebro y el corazón pueden caracterizar el estado de conciencia de los pacientes, y (2) determinar si la información extraída de la actividad cardiaca (EKG) complementa la información obtenida del EEG. Las hipótesis de trabajo son: (1) los marcadores de actividad cardíaca como la frecuencia y la variabilidad van a presentar los mismos resultados que en trabajos anteriores y (2) marcadores de modulación de la actividad cardíaca medidas bajo el test Local-Global (ver 1.5.2) van a estar presentes en las mismas condiciones que las respuestas neuronales medidas con EEG. En este último caso, se espera que el efecto Local no distinga los grupos de pacientes (Faugeras et al., 2011), y el efecto Global va a estar presente sólo en pacientes con un nivel de conciencia preservado (Faugeras et al., 2012; King et al., 2013a).

Para probar estas hipótesis, utilizamos un grupo de 259 pacientes registrados con EEG y el paradigma Local-Global. Sin embargo, estos registros no presentaban EKG (electrocardiograma). Utilizando la técnica de descomposición de la información mutua (Bell and Sejnowski, 1995; Lee et al., 1999; Hyvärinen and Oja, 2000), extrajimos las series temporales correspondientes a la actividad cardiaca en 132 pacientes. Una vez obtenidos, utilizamos un algoritmo de detección de picos para obtener las series temporales de los latidos del corazón, calculando la frecuencia cardíaca, su variabilidad, y la ubicación de los latidos con respecto a los estímulos auditivos del paradigma Local-Global.

La actividad cardiaca de base (frecuencia y variabilidad), no mostró diferencias entre los grupos de pacientes MCS y VS/UWS. Sin embargo, encontramos una correlación positiva entre el valor de CRS-R y los marcadores de frecuencia y variabilidad, aunque sólo para el grupo de pacientes VS/UWS (Figura 4.4). Creemos que esto se relaciona con el estado físico de los pacientes, al presentar condiciones de taquicardia en pacientes con CRS-R bajos.

En cuanto a las modulaciones de la actividad cardiaca por el paradigma auditivo, realizamos el mismo contraste de efectos locales y globales, pero midiendo las diferencias de tiempo entre el estímulo auditivo los latidos anterior (PRE) y posterior (POST).
4.6. DISCUSSION

Cuando contrastamos los intervalos PRE (latidos previos al estimulo), no encontramos ninguna diferencia entre los grupos VS/UWS y MCS tanto para el efecto Local como para el efecto Global. Sin embargo, los intervalos POST presentaron una diferencia sólo en el grupo MCS y sólo en el efecto Global (Figura 4.7). El resultado fue un acortamiento del intervalo entre el estimulo y el latido posterior cuando el estímulo correspondía con la tarea a realizar.

Para determinar si esta modulación de la actividad cardíaca es o no redundante con respecto a la información obtenida del EEG, entrenamos un clasificador SVM con combinaciones de 3 distintos grupos de features: (1) Marcadores de EEG (ver 2.5), (2) marcadores de actividad cardíaca de base (frecuencia y variabilidad) y (3) marcadores de procesos cognitivos en el EKG (intervalos PRE y POST para los contrastes Locales y Globales). Este análisis demostró un incremento significativo en la performance cuando los marcadores de procesos cognitivos de EKG fueron incluidos (Figura 4.8).

En conclusión, mostramos una relación entre la función del sistema nervioso autónomo y estímulos auditivos que se observa sólo en situaciones con un nivel preservado de conciencia. Nuestros resultados sugieren que la modulación de la actividad cardíaca es relevante para el diagnóstico de pacientes DOC, y que la información obtenida de esta modulación es parcialmente independiente de la información neuronal. Este trabajo sienta la bases para el estudio de los pacientes DOC desde el paradigma incorporado, en el cual las interacciones entre el cuerpo y el cerebro contribuyen al enfoque holístico del procesamiento consciente.
Chapter 5

Summary and final remarks

In chapter one, we described the fundamentals of EEG, the problematic of DOC, and the current state-of-the-art tools for the diagnosis of DOC. In chapter two, we analysed the validity and robustness of an automatic processing and DOC diagnostic method based on EEG-extracted markers. We tested this method under distinct simulated and real conditions from different clinical centres, concluding that the model extracts reliable signatures of consciousness, maintaining the diagnostic accuracy across conditions. In chapter three, we aimed at extending our knowledge on multisensory integration and states of consciousness. For this purpose, we introduced a modification of an auditory paradigm, currently used to assess the diagnosis of DOC patients, to include somatosensory and visual stimulation. We tested this paradigm on healthy controls, obtaining results consistent with previous works on multisensory integration. In chapter four, we moved away from a strict neurocentric approach for the study of consciousness and included bodily signals in our analysis. We found that the already known brain-heart interactions are mediated by the state of consciousness. Interestingly, this modulation of cardiac activity by cognitive processes is present only in the group of DOC patients showing high-order behavioural responses.

The present work exceed the current state of knowledge at two dimensions. Clinically, our work on automated EEG-based diagnosis confirms and expands the utility of machine learning as an assisting technology for the clinical management of DOC. On the one hand, the here proposed automated diagnostic tool validates previous findings. On the other, it generalises to other experimental setups and recording conditions. Therefore, our results provide support for the use of EEG-extracted markers and machine learning in clinical settings, in a flexible and reliable way. At the same time, it permits the extrapolation to other experimental application, such as the multisensory cross-modal investigation as a supplementary marker of conscious state. In that case, the here proposed methods can be used synergically with other physiologically-extracted
information.

Scientifically, it sheds light on the relationship between states of consciousness and the embodied paradigm to cognition. To date, consciousness is directly inferred by means of subjective reports, task performance and by observing nonreflex behaviours. In the absence of subjective reportability, like in DOC patients, it is challenging to know whether patients retain any form of subjective experiences. According to cognitive science, subjectivity is a construct which can be approached hierarchically, from the experience of a "me" shaped by perceptions of others and their perceptions of me (social self), to being a continuous person over time (narrative self), to the experience of a sense of reality of the world and of me within it (minimal self) (Seth, 2013). Especially the case of the minimal self can be understood as the conscious experience of being someone, which is pre-reflective in nature, i.e. independent from explicit cognition and linguistic properties (Gallagher, 2000). Contemporary neurocognitive approaches, which are formulated based on reportable experiences, imply that undifferentiated brain activity might account for the inability to retain subjective experience. Therefore, the self in unconscious conditions is severely compromised and therefore absent. Alternatively, embodiment, a position in cognitive neuroscience and philosophy of mind, provides a more specific framework for the study of minimal selfhood by emphasising the role of body in shaping cognition (Varela et al., 1991). According to the theory, an organism is considered to be a self when the following conditions are jointly met: a) it possesses volume in space (localised within bodily boundaries), b) it recognises a global body representation (the body is perceived as a whole as opposed to localised body parts and isolated movements), and c) it possesses a visuospatial frame of reference/egocentric model of reality. In other words, subjectivity in its fundamental form is the process of the conscious experience of being a distinct, holistic entity, embodied and embedded in space and time (Blanke and Metzinger, 2009). Consequently, to infer the presence of a minimal self “it is sufficient to show a passive, multisensory and globalised availability of an integrated, transparent and global representation of the spatiotemporally situated body” (Blanke and Metzinger, 2009). The here presented work on brain-heart interaction bridges the empirical gap of the relationship between subjective experience and altered states of consciousness in DOC because it extends from a merely neurocentric approach. It shows that human cognition is realised holistically adding to growing evidence that the mind is a dynamic process between the organism and the world.
Resumen en español

En el capítulo uno, describimos los fundamentos del EEG, la problemática de los desordenes de conciencia y las herramientas actuales para el diagnóstico de DOC. En el capítulo dos, analizamos la validez y robustez de una herramienta automatizada para el diagnóstico de DOC basado en la extracción de marcadores de EEG. Probamos este método bajo diversas situaciones simuladas y reales con datos de distintos centros clínicos, concluyendo que el modelo extrae marcadores confiables de conciencia, manteniendo la precisión en el diagnóstico. En el capítulo 3, apuntamos a extender el conocimiento con respecto a la integración multisensorial y los estados de conciencia. Para ello, introducimos una modificación a un paradigma de estimulación auditivo usado en la actualidad para el diagnóstico de DOC, agregando estímulos somatosensoriales y visuales. Probamos este paradigma en sujetos sanos, obteniendo resultados consistentes con los trabajos previos en integración multisensorial. En el capítulo 4, nos movemos del enfoque estrictamente neurocéntrico para el estudio de la conciencia e incluimos señales corporales en nuestro análisis. Encontramos que los procesos cognitivos producen una modulación de la actividad cardíaca, que se presenta sólo en el grupo de pacientes que tienen respuestas comportamentales mas complejas.

Este trabajo excede los conocimientos actuales en dos dimensiones. Clínicamente, nuestro trabajo en el diagnóstico automatizado de DOC, confirma y expande la utilidad del aprendizaje automático como una herramienta para el manejo clínico de pacientes DOC. Por un lado, esta herramienta valida los resultados obtenidos anteriormente. Por el otro, muestra que se puede generalizar a otras condiciones experimentales y de adquisición. Por lo tanto, nuestros resultados proveen evidencia para el uso de marcadores extraídos de EEG y aprendizaje automático en aplicaciones clínicas de manera flexible y confiable. Al mismo tiempo, permite la extrapolación a otras aplicaciones experimentales, como la investigación de nuevos marcadores de conciencia basados en integración multisensorial y cроссsensorial. En ese caso, los marcadores aquí propuestos se pueden utilizar sinérgicamente con otra información fisiológica.

Científicamente, aclara la relación entre estados de conciencia y el paradigma incorporado para la cognición. Hasta ahora, la conciencia es directamente inferida por medio de reportes subjetivos, la práctica de tareas y la observación de comportamiento no-reflexivo. En la ausencia de reportabilidad subjetiva, como en los pacientes DOC, resulta desafiante saber si estos pacientes retienen alguna forma de experiencia subjetiva. De acuerdo a la ciencia cognitiva, la subjetividad es una construcción que puede ser abordada jerárquicamente, desde la experiencia de un "yo" (individuo) moldeado por percepciones de otros y de sus percepciones de uno mismo (un individuo social), a ser una persona continua en el tiempo (un individuo narrativo), a la experiencia de un sentido de realidad del mundo y el individuo dentro de el (un individuo minimal).
(Seth, 2013). Especialmente, el caso del individuo minimal puede ser comprendido como la experiencia consciente de ser alguien, quien es prereflectivo en su naturaleza, i.e. independiente de cognición explícita y propiedades lingüísticas (Gallagher, 2000). Enfoques neurocognitivos contemporáneos, cuya formulación se basa en experiencias reportables, implican que la actividad cerebral indiferenciable podría explicar la incapacidad de retener una experiencia subjetiva. Por lo tanto, el individuo en condiciones inconscientes se ve severamente comprometido y, en consecuencia, ausente. Alternativamente, *embodiment* (del paradigma incorporado), es una posición en neurociencias cognitivas y filosofía de la mente que provee un marco de referencia más específico para el estudio de la individualidad minimal, enfatizando el rol del cuerpo en la formación de la cognición (Varela et al., 1991). Según la teoría, un organismo es considerado un individuo cuando se cumplen las siguientes condiciones en forma conjunta: a) posee volumen en el espacio (localizado dentro de límites corporales), b) reconoce un cuerpo global y c) posee, en forma visuoespacial, un marco de referencia o modelo egocéntrico de la realidad. En otras palabras, la subjetividad en su forma fundamental es el proceso de la experiencia consciente de ser una entidad holística distinta, incorporada y embueida en el espacio y el tiempo (Blanke and Metzinger, 2009). En consecuencia, para inferir la presencia de un individuo minimal, “es suficiente mostrar una disponibilidad pasiva, multisensorial y globalizada, de una representación del cuerpo situado en el espacio y el tiempo, de manera integrada, transparente y global” (Blanke and Metzinger, 2009). El trabajo presentado aquí en interacciones cerebro-corazón achica la brecha empírica de la relación entre la experiencia subjetiva y los estados de conciencia alterados en pacientes DOC, porque lo aparte de un enfoqué primordialmente neurocéntrico. Muestra que la cognición humana se construye holísticamente, indicando que la mente es un proceso dinámico entre el organismo y el mundo.
Bibliography


Tim Bayne, Jakob Hohwy, and Adrian M. Owen. Are There Levels of Consciousness?, 2016. ISSN 1879307X. 2, 23
Henry Beecher, Raymond Adams, A Clifford Barger, William Curran, Derek Denny-Brown, Dana Farnsworth, Jordi Folch-Pi, Everett Mendelsohn, John Merrill, Joseph Murray, Ralph Potter, Robert Schwab, and William Sweet. A definition of irreversible coma: report of the ad hoc committee of the Harvard Medical School to examine the definition of brain death, 1968. ISSN 0041-1337. 3, 23


James L. Bernat. Ethical issues in the perioperative management of neurologic patients, 2004. ISSN 07338619. 3


Frédéric Faugeras, Benjamin Rohaut, Mélanie Valente, Jacobo Sitt, Sophie Demeret, Francis Bolgert, Nicolas Weiss, Alexandra Grinea, Clémence Marois, Marion Quirins, Athena Demertzi, Federico Raimondo, Damien Galanaud, Marie-Odile Habert, Denis Engemann, Louis Puybasset, and Lionel Naccache. Survival and consciousness recovery are better in the minimally conscious state than in the vegetative state. *Brain Injury*, pages 1–6, nov 2017. ISSN 0269-9052. doi: 10.1080/02699052.2017.1364421. 3


Thomas Frodl-Bauch, Ronald Bottlender, and Ulrich Hegerl. Neurochemical substrates and neuroanatomical generators of the event-related P300, 1999. ISSN 0302282X. 13, 14


Joseph T. Giacino, Kathleen Kalmar, and John Whyte. The JFK Coma Recovery Scale-Revised: Measurement characteristics and diagnostic utility, 2004. ISSN 00039993. 5, 17, 23


Guy Kahane and Julian Savulescu. Brain damage and the moral significance of consciousness, 2009. ISSN 03605310.


J. R. King and S. Dehaene. Characterizing the dynamics of mental representations: The temporal generalization method, 2014. ISSN 1879307X.


Steven Laureys. Death, unconsciousness and the brain, 2005a. ISSN 1471003X. 6


Steven Laureys and Melanie Boly. What is it like to be vegetative or minimally conscious? *Current opinion in neurology*, 20(6):609–613, 2007. ISSN 1350-7540. doi: 10.1097/WCO.0b013e3282f1d6dd. 2


Gilles Louppe. Understanding Random Forests: From Theory to Practice. jul 2014. 49


Charles-Edouard Luyt, Damien Galanaud, Vincent Perlberg, Audrey Vanhaudenhuyse, Robert D Stevens, Rajiv Gupta, Hortense Besancenot, Alexandre Krainik, Gérard Audibert, Alain Combes, Jean Chastre, Habib Benali, Steven Laureys, and Louis


Sepideh Sadaghiani and Andreas Kleinschmidt. Brain Networks and α-Oscillations: Structural and Functional Foundations of Cognitive Control, 2016. ISSN 1879307X.


Nicholas D. Schiff. Recovery of consciousness after brain injury: a mesocircuit hypothesis, 2010. ISSN 01662236. 50


Caroline Schnakers, Joseph Giacino, Kathleen Kalmar, Sonia Pitet, Eduardo Lopez, Mélanie Boly, Richard Malone, and Steven Laureys. Does the FOUR score correctly diagnose the vegetative and minimally conscious states? [1], 2006. ISSN 03645134. 5


Pasquale Striano, Francesca Bifulco, and Giuseppe Servillo. The saga of Eluana Englaro: Another tragedy feeding the media, 2009. ISSN 03424642. 3


Giulio Tononi and Christof Koch. The neural correlates of consciousness: An update, 2008. ISSN 00778923. 16


