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Coupled Correlates of Attention and Consciousness

by
Ravi Chacko

A dissertation presented to
The Graduate School
of Washington University
in partial fulfillment of the
requirements for the degree
of Doctor of Philosophy

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Table of Contents

List of Figures	iv
List of Tables	iv
Preface	v
Acknowledgments.....	vi
Abstract.....	viii
1) Introduction: Identifying Electrophysiological Correlates of Attention and Consciousness.....	1
1.1 Brain Computer Interfaces (BCIs)	1
1.2 The Problem: BCI for Hemineglect:	3
1.3 The Question: How are attention and consciousness represented in the brain?.....	4
2) Background: Attention and Consciousness	8
2.1 Attention and inattention.....	8
2.1.1 Hemineglect (HN).....	9
2.1.2 The Neuroanatomy of (In)Attention:	10
2.1.3 Hemineglect Rehabilitation and a Rationale for BCI	12
2.1.4 Measuring Attention: The Posner Task.....	13
2.1.5 The Electrophysiology of Attention.....	15
2.2 Consciousness	19
2.2.1 Defining Consciousness.....	19
2.2.2 Sleep.....	21
2.2.3 Propofol Induced Loss of Consciousness (PILOC)	23
3) Methods: Measuring Phase-Amplitude Coupling.....	25
3.1 From Neuronal Firing to Scalp Recordings	25
3.2 Phase-amplitude coupling (PAC).....	30
3.3 Measuring PAC.....	31
3.4 Theory of PAC in neural oscillations.....	34
3.5 Challenges to PAC theory and measurement: Non-sinusoidal, Sharp Waves	35
3.6 Evidence for Phase-amplitude coupling in Attention and Consciousness	36

4) Distinct Phase-Amplitude Couplings Distinguish Cognitive Processes in Human Attention	38
4.1 Abstract	38
4.2 Introduction.....	39
4.3 Materials and Methods.....	41
4.4 Results.....	47
4.5 Discussion	61
4.6 Supplemental Figures:	65
5) Alpha-phase Coupling Distinguishes Mechanistically Distinct Unconscious States	68
5.1 Introduction.....	68
5.2 Results.....	71
5.3 Discussion	76
5.4 Supplemental Data	78
6) Proof of Concept and Conclusion.....	84
Appendix A: A Review of Rehabilitation Paradigms for Hemineglect.....	90
References.....	92

List of Figures

Title	Page
3.1.1 Signal processing and analysis methods	38
3.1.2 Amplitude and phase of a complex signal	39
3.2.1 Phase-amplitude coupling measurement with modulation index	42
3.3.1 Importance of high frequency-for-amplitude wavelet bandwidth	43
4.4.1 Task design showing analysis periods and behavioral results	57
4.4.2 Identification of distinct phase-amplitude coupling clusters	59
4.4.3 Functional classification of “spatial” and “behavioral” sites	62
4.4.4 Phase-preferences and relative PAC magnitude distinguishes frequency and class	64
4.4.5 PAC magnitude differences across task conditions	67
4.4.6 Transient waves are behaviorally relevant and coupled to 7.2 Hz phase	69
4.6.1 Supplemental figures	75
5.2.1 Double dissociation of PAC by frequency pair and sleep state	81
5.2.2 TABL PAC decreases during SWS but increases during PLOC	83
5.2.3 Increased theta- and alpha-phase PAC is not caused by changes in power	85
5.4.1 Consistency of double dissociation across patients	88
5.4.2 Delta-phase PAC increases and alpha-phase PAC decreases with NREM stage	92
6.1.1 Log-loss classifier performance for left vs right target locations	95
6.1.2 Adding a dimension to the spectrum of consciousness	99

List of Tables

Table 1: NIH stroke scale for quantification of neglect	18
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Preface

It has been an honor and a privilege to study the brain at Washington University in St. Louis. I have been fortunate to work with researchers across many disciplines, such as biomedical engineering, experimental and theoretical neuroscience, cognitive neuropsychology, neurosurgery, neurology, neuronal modelling, anesthesiology, hypnology, machine learning, probability, signal processing and computer science. The following work will touch on high-level concepts from all these fields to describe electrophysiological correlates of attention and consciousness. Chapter 1 supplies the motivation for our thesis and includes a rationale for the topics discussed. The goal of this project was to better understand the mesoscopic electrophysiology of attention so that it can be used in the rehabilitation of attentional disorders such as hemispatial neglect. This goal led to a study of phase-amplitude coupling and its relationship to attention and arousal. Chapter 2 is a background on attention, sleep and anesthesia. Chapter 3 reviews the methodological choices we make for measuring phase-amplitude coupling (PAC). Chapter 4 reports our experimental findings on how different PAC frequency clusters relate to different aspects of human attention. Chapter 5 reports findings on differences in PAC between conscious and two distinct unconscious states. We conclude with a discussion of how these findings can be used in future work to implement brain computer interfaces for rehabilitating hemispatial neglect in Chapter 6.

Acknowledgments

I am grateful for the many discussions with my fellow graduate students, including Nick Szrama, Mrinal Pahwa, Carl Hacker, Ammar Hawasli, Jarod Roland, Mohit Sharma, David Bundy, Joey Humphries, Andy Daniels, DoHyun Kim, Josh Siegel and Amy Daitch. Without them this research would never have come to fruition. I greatly appreciate the efforts of my mentor, Eric Leuthardt, pushing me to consider new research topics. I appreciated the complementary mentorship of Maurizio Corbetta and Gordon Shulman, especially when discussing meaning of our experimental findings. I also enjoyed discussing the theoretical underpinnings of our modelling work and what the definition of “is” is with ShiNung Ching. I am grateful to the additional mentorship afforded to me by Linda Larson-Prior, Barani Raman, Dan Moran and Andy Mitz, who all pushed me to be a better scientist and engineer. I consider the time spent bantering about theoretical neuroscience priceless. I especially enjoyed questioning the foundational premises of my work, and of many other people’s work, as it gave me a great deal of perspective. Finally, I must thank my father, for coaching me through a PhD, my mother, for understanding me, my brother, for being my inspiration and my fiancé, for encouraging me through the tough times while accomplishing her own incredible feats.

Ravi Chacko

Washington University

May 2019

This work is dedicated to my dear parents.

Abstract

ABSTRACT OF THE DISSERTATION

Coupled Correlates of Attention and Consciousness
by

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Washington University in St. Louis, 2019

Eric C. Leuthardt, Chair

Introduction: Brain Computer Interfaces (BCIs) have been shown to restore lost motor function that occurs in stroke using electrophysiological signals. However, little evidence exists for the use of BCIs to restore non-motor stroke deficits, such as the attention deficits seen in hemineglect. Attention is a cognitive function that selects objects or ideas for further neural processing, presumably to facilitate optimal behavior. Developing BCIs for attention is different from developing motor BCIs because attention networks in the brain are more distributed and associative than motor networks. For example, hemineglect patients have reduced levels of arousal, which exacerbates their attentional deficits. More generally, attention is a state of high arousal and salient conscious experience. Current models of consciousness suggest that both slow wave sleep and Propofol-induced unconsciousness lie at one end of the consciousness spectrum, while attentive states lie at the other end. Accordingly, investigating the electrophysiology underlying attention and the extremes of consciousness will further the development of attentional BCIs.

Phase amplitude coupling (PAC) of neural oscillations has been suggested as a mechanism for organizing local and global brain activity across regions. While evidence suggests that delta-high-gamma PAC, which includes very low frequencies (i.e. delta, 1-3 Hz) coupled with very high frequencies (i.e. gamma 70-150 Hz), is implicated in attention, less evidence exists for the involvement of coupled mid-range frequencies (i.e. theta, 4-7Hz, alpha: 8-15 Hz, beta: 15-30 Hz and low-gamma: 30-50 Hz, aka TABL PAC). We found that TABL PAC correlates with reaction time in an attention task. These mid-range frequencies are important because they can be used in non-invasive electroencephalography (EEG) BCI's. Therefore, we investigated the origins of these mid-frequency interactions in both attention and consciousness. In this work, we evaluate the relationship between PAC to attention and arousal, with emphasis on developing control signals for an attentional BCI.

Objective: To understand how PAC facilitates attention and arousal for building BCI's that restore lost attentional function. More generally, our objective was to discover and understand potential control features for BCIs that enhance attention and conscious experience.

Methods: We used four electrophysiological datasets in human subjects. The first dataset included six subjects with invasive ECoG recordings while subjects engaged in a Posner cued spatial attention task. The second dataset included five subjects with ECoG recordings during sleep and awake states. The third dataset included 6 subjects with invasively monitored ECoG during induction and emergence from Propofol anesthesia. We validated findings from the second dataset with an EEG dataset that included 39 subjects with EEG and sleep scoring.

We developed custom, wavelet-based, signal processing algorithms designed to optimally calculate differences in mid-frequency-range (i.e. TABL) PAC and compare them to DH PAC across different attentional and conscious states. We developed non-parametric cluster-based

permutation tests to infer statistical significance while minimizing the false-positive rate. In the attention experiment, we used the location of cued spatial stimuli and reaction time (RT) as markers of attention. We defined stimulus-related and behaviorally-related cortical sites and compared their relative PAC magnitudes. In the sleep dataset, we compared PAC across sleep states (e.g. Wake vs Slow Wave Sleep). In the anesthesia dataset, we compared the beginning and ending of induction and emergence (e.g. Wake vs Propofol Induced Loss of Consciousness)

Results: We found different patterns of activity represented by TABL PAC and DH PAC in both attention and sleep datasets. First, during a spatial attention task TABL PAC robustly predicted whether a subject would respond quickly or slowly. TABL PAC maintained a consistent phase-preference across all cortical sites and was strongest in behaviorally-relevant cortical sites. In contrast, DH PAC represented the location of attention in spatially-relevant cortical sites.

Furthermore, we discovered that sharp waves caused TABL PAC. These sharp waves appeared to be transient beta (50ms) waves that occurred at ~140 ms intervals, corresponding to a theta oscillation. In the arousal dataset DH PAC increased in both slow wave sleep (SWS) and Propofol-induced loss of consciousness (PILOC) states. However, TABL PAC increased only during PILOC and decreased during SWS, when compared to waking states. We provide evidence that TABL PAC represents “gating by inhibition” in the human brain.

Conclusions: Our goal was to develop electrophysiological signals representing attention and to understand how these features explain the relationship between attention and low-arousal states.

We found a novel biomarker, TABL PAC, that predicted non-spatial aspects of attention and discriminated between two states of unconsciousness. The evidence suggested that TABL PAC represents inhibitory activity that filters out irrelevant information in attention tasks. This inhibitory mechanism of was confirmed by significant increases in TABL PAC during Propofol

anesthesia, when compared to SWS or waking brain activity. We conclude that TABL PAC informs the development of electrophysiological control signals for attention and the discrimination of unconscious states.

1) Introduction: Identifying Electrophysiological Correlates of Attention and Consciousness

“Every speculative enterprise which he undertook, and they were many and various, was carried to sure success by the same qualities of cool, unerring judgment, far-reaching sagacity, and apparently superhuman power of organizing, combining, and controlling, which had made him in politics the phenomenon of the age” – *The Ablest Man in the World*, Edward Page Mitchell, 1884

In Edward Page Mitchell’s short story, “The Ablest Man in the World” a doctor describes treating a sick Russian baron and stumbling on a curious finding. Underneath the baron’s skull cap, a silver dome covered a clockwork brain. The baron was born with neurological deficits and a Russian doctor with skill in watchmaking designed a clockwork brain to fix them. Not only did it fix his deficits, but it gave him superhuman powers. For more than a century human have imagined using devices to improve neurological function. This work extends this effort by understanding how the brain represents attention and certain aspects of consciousness to treat disease and augment human cognition.

1.1 Brain Computer Interfaces (BCIs)

The purpose of a BCI is to decode the neural activity associated with a behavior so that the behavior can be recreated or augmented. BCIs are designed by modelling and predicting brain activity with high accuracy in over short time periods. To do this, we typically record brain activity while subjects repeat a task. We then model the neural activity that represents important

aspects of the task and use that model to predict behavior. For example, a paralyzed patient with an injured spinal cord has a functioning motor cortex. We might ask the patient to think about moving their hand while recording brain activity from their motor cortex. After recording enough data, a model can be created to predict when the patient is thinking about moving his hand. Ultimately a robotic hand can complete the required behavior after the model makes an appropriate prediction, thus recreating the patient's lost neurological function. BCIs typically consist of three elements, 1) a neural interface, 2) a computational processing module and 3) an output. While BCIs have made progress in the restoration of motor deficits in stroke and paralysis¹⁻⁴, they have not succeeded in improving cognitive functions like the ones in Mitchell's quote above. Here we focus on the cognitive function called attention. Developing BCIs to augment attention would benefit stroke patients with Hemineglect.

Hemineglect is an attention disorder and the motivation for this thesis. Previously, BCIs have been used to improve hemiparetic stroke syndromes, which incorporate movement disorders¹. When we attempted to apply BCI principles from movement disorders to attention disorders several issues became salient. Compared to movement, attention is a higher order cognitive function that involves more brain networks working in concert. For example, if you call out to someone and they don't respond, it could be because (A) they don't perceive you calling them (e.g. they're deaf), (B) they don't have the ability to execute a response (e.g. they're paralyzed) or (C) there is something wrong with their brain's ability to convert a perception of you calling them to a response (e.g. Hemispatial neglect). Our goal was to understand the electrophysiological correlates of attention, which transforms sensory perceptions into measurable behaviors, so that they can be used to repair neurological deficits. The following two

chapters discuss how a clinical and engineering motivation required the scientific approach we pursued.

1.2 Problem: BCI for Hemineglect:

Hemineglect (HN) describes the inability to attend to one half of space (i.e. hemifield) that typically follows a right-brain stroke. These patients fail to shave one half of their face, dress one half of their body and draw one half of pictures they are given to copy. They also have slower response speeds to stimuli in both hemifields and suffer from reduced levels of arousal. HN occurs in 25-30% of all stroke patients, 10-13% of left hemisphere strokes and 40-82% of right hemisphere strokes⁵⁻⁹. This amounts to roughly 200,000 people a year. Decades of research have yielded theories that explain elements of HN¹⁰⁻¹³, but meta-analyses conclude that there currently is no effective rehabilitation treatments for HN^{14,15}. The goal of this work was to develop innovations in HN rehabilitation using BCI methodologies. If we can understand which signals explain lateralized attention and response speed, we can teach HN patients to increase their output of that signal. To develop these BCIs we proposed the development of attentional control signals, or electrophysiological signatures of attention.

We focus on electrophysiological signatures because they can be used with electroencephalography (EEG) neurofeedback. EEG uses non-invasive scalp electrodes that record electric potential. While an EEG neurofeedback device is the ultimate goal, this research focuses on electrocorticography (ECoG), which uses surgically implanted electrodes on the surface of the brain. ECoG is prescribed for epilepsy patients, who have failed conservative treatment, in order to localize the origins of seizures. The advantage of ECoG is that the signal is less noisy and more spatially specific than EEG signal. ECoG also records higher frequencies

than EEG. However, our goal will be to focus on features that can eventually be used with EEG feedback, which rules out frequencies above 50 Hz.

Finally, ideal attentional control features will be measurable on single trials. If an EEG neurofeedback paradigm for hemineglect mirrors its motor stroke counterpart, then it will require the patient to practice. This means many repeated trials. The neurofeedback device must tell its operator how well he or she attended during the last trial. If we cannot compute this value quickly, then the operator won't know how to improve their attention. This is a consequence of operant conditioning, where the strength of a behavior is modified by reward or punishment (i.e. feedback).

In summary, our goal is to develop attentional control features that will serve an EEG neurofeedback device. It must use frequencies below 50 Hz and must be able to detect whether the subject paid attention on a single trial. In the end, a hemineglect patient will wear an EEG headset while playing a computer game that challenges their attention. On trials where they attended properly, they will be rewarded. On trials where they fail to attend, they will not be rewarded. By repeatedly playing this game, they will learn what promotes and prevents attention. But what does it mean to attend properly? Hemineglect patients have more than one deficit. Not only are they unable to attend to one half of space as well as the other, they also have deficits that exist on both halves of space^{16,17}. Furthermore, the colloquial usage of the word "attention" doesn't always fit the scientific and clinical understanding. Therefore, before we set out to find attentional control signals, we will first understand the components of attention.

1.3 The Question: How are attention and consciousness represented in the brain?

To begin to understand the lateralized attention deficit in hemineglect, think of your attention like a spotlight. Much of the time it moves around with your gaze. However, just as your eyes

can scan words on a page without absorbing any of its meaning, your attention can move independently of your gaze. To convince yourself, fixate on the red dot below after completing this passage and avoid looking directly at the items to the left or the right of the dot. Try attending to the left of the dot without looking away from the dot. You'll notice that item to the left of the page will become more salient and you will make out its identity more easily. Notice how shifting your attention to the left of the dot (again without breaking fixation) makes it more difficult to resolve the stimulus to the right of the page.



This is because attention is a limited resource, or a “limited-capacity spotlight”¹⁸. When it shines in one part of the visual field, it does not shine in another. HN patients fail to attend to one half of space. This is known as the “spatial” deficit in HN. EEG correlates of lateralized attentional shifts have been shown in the alpha (8-13 Hz) frequency¹⁹. However, alpha power did not discriminate left and right shifts in attention much better than chance. Furthermore, lateralized attention deficits aren’t the only attentional deficits in hemineglect.

In healthy subjects, attention is often measured using response times (RT)²⁰⁻²². A subject typically attends to a location in space, then respond to stimuli that appear at that location or at an unattended location. RT measures the entire mental sequence of deploying attention, orienting it, sustaining it, sensing or perceiving the relevant stimuli, processing it, then planning and executing a motor response. In addition to the markedly slower responses (i.e. higher RTs) to stimuli in their neglected hemifield (i.e. the “spatial” deficit), HN patients also have a slower response to all stimuli, regardless of where it occurs²³. This is known as the “non-spatial” deficit in neglect. What’s more, depending on the location of the stroke, an HN patient may have different ratios of “spatial” and “non-spatial” deficits in neglect²⁴. Therefore, we must be careful to distinguish between “spatial” and “non-spatial” aspects of attention. But “non-spatial” attention is not the only cognitive function to influence RT. For example, when you are drowsy you will respond more slowly to all stimuli. Furthermore, if you are distracted by some other task, you will also respond more slowly to all stimuli. Therefore, we must understand how attention is related to consciousness more generally.

Consciousness refers to a multifaceted concept, but we will initially describe it with two dimensions. The first is arousal or wakefulness, the second is awareness or experience²⁵. In clinical settings, the first dimension is useful because distinguishes levels of brain injury or coma. The second dimension is the subject of philosophical writings and psychophysics research. Most of the cognitive states discussed below differ along both axes of arousal and awareness. For example, a state of attention is a state of high arousal and high awareness compared to resting wakefulness. Conversely, sleep is a state of low arousal and low awareness. In the following chapters, we will compare what we learned from attentive states to two states of lost consciousness, slow-wave sleep (SWS) and Propofol-induced loss of consciousness

(PILOC). We use these two states because SWS has a well-studied neurophysiology and PILOC has a well-known mechanism of action.

In summary, our engineering goal of developing a BCI for Hemineglect requires the scientific pursuit of measuring physiology associated with the abstract concepts of attention and consciousness. Therefore, we will study attention as it relates to a) lateralized shifts in attention and b) reaction time (RT), with the caveat that RT are generated by multiple cognitive systems. Furthermore, we will investigate consciousness across two states, SWS and PILOC. Contrasting brain activity across these variables and states underlies our scientific findings.

2) Background: Attention and Consciousness

Attention is a cognitive¹ selection mechanism that selects an object or idea for further processing. A colloquial example of inattention is when one attempts to read words on a page and realizes, at the end of a passage, that the passage wasn't read or comprehended. Even though eyes scanned words on the page, the cognitive function called attention did not select the words for language comprehension. Thus, the object (i.e. words) that one's eyes selected was not selected by one's brain for further processing (i.e. comprehension). With respect to spatial attention, where you look is commonly where you attend. But as the example demonstrates, this is not always the case. A behavioral paradigm that measured this type of inattention is the Posner cueing task²⁶. As this task is central to the study described in Chapter 4, we will use it to elaborate neuropsychological concepts around attention and clinical sequelae of lost attentional function in hemineglect (HN).

2.1 Attention and inattention

William James's famously said "[Attention] is the taking possession by the mind, in clear and vivid form, of one out of what seem several simultaneously possible objects or trains of thought, localization, concentration, of consciousness are of its essence." Simply put, attention is a cognitive function that selects what the brain thinks about. This description, however, is lacking because we have an incomplete understanding of how the brain processes "items or trains of thought" that have been selected. Attention studies typically measure behaviors as a

¹ Cognition is the mental action or process of acquiring knowledge and understanding through thought, experience, and the senses. Or a result of this; a perception, sensation, notion, or intuition.

proxy for cognitive processing. An experiment will manipulate inputs (e.g. items to be selected) and measure outputs (e.g. actions required to complete the task). We therefore study how brain transforms sensory inputs into motor outputs. Unfortunately, this “black box” model of attention is also vague. Therefore, we will motivate our definition of attention by two HN deficits that are measurable with the Posner spatial cueing task. The first deficit is in spatially lateralized aspects of attention, the second deficit is in non-spatial aspects of attention.

2.1.1 Hemineglect (HN)

Hemineglect (a.k.a. neglect, hemispacial neglect and spatial neglect) is a stroke syndrome that is most commonly diagnosed by a lateralized deficit in attention. HN is diagnosed as hemi-inattention (**Table 1**), a failure to orient, or respond to stimuli on one side of space that cannot be explained by a visual or motor deficit^{8,27}. Typically, right sided parietal strokes lead to a left sided hemineglect. A patient with profound hemineglect may orient their head to the right of center and may not use their left hand.

<p>11. Extinction and Inattention (formerly Neglect): Sufficient information to identify neglect may be obtained during the prior testing. If the patient has a severe visual loss preventing visual double simultaneous stimulation, and the cutaneous stimuli are normal, the score is normal. If the patient has aphasia but does appear to attend to both sides, the score is normal. The presence of visual spatial neglect or anosagnosia may also be taken as evidence of abnormality. Since the abnormality is scored only if present, the item is never untestable.</p>	<p>0 = No abnormality.</p> <p>1 = Visual, tactile, auditory, spatial, or personal inattention or extinction to bilateral simultaneous stimulation in one of the sensory modalities.</p> <p>2 = Profound hemi-inattention or extinction to more than one modality; does not recognize own hand or orients to only one side of space.</p>
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Table 1: NIH Stroke scale for the quantification of (Hemi)neglect. Extinction is when hemineglect symptoms occur only when stimuli are presented simultaneously in both hemifields.

The “spatial” or “lateralized” deficits in neglect are most striking. HN patients may shave only one half of their face or dress only one side of their body. It’s important to note that HN is not a visual or tactile deficit. Patients can see into the neglected hemispace and feel their

neglected hemibody when cued to do so. Furthermore, it is not a motor problem, as patients can move their limbs and respond to directions. Therefore, it is a problem with attention networks in the brain that coordinate sensory inputs with motor outputs to execute goal-oriented behaviors. Patients with HN may not feel anything is wrong. It is as if their world has shrunk to half the size, but this shrunken world has stretched to occupy the entire space. What's more, HN doesn't only affect present experiences, it also interacts with memory. One hemineglect patient was asked to imagine standing in their town square and recall all the buildings he could. He dutifully recalled all the buildings on the right of where he was standing. Next, the clinician asked this patient to turn around, in his imagination, and repeat the task. Again, the patient recalled all the buildings to his right. Without realizing it, the patient had recounted all the buildings in the square, but at any given positioning he could only recall one side. This points to the multifaceted nature of attention, beyond its effects on stimulus perception and motor behavior²³.

“Non-spatial” HN deficits are not lateralized to one side. HN patients are slower to respond to any stimuli, regardless of the side they are stimulated on. They also suffer deficits in sustained attention and working memory. These deficits have been shown to predict clinical outcome independently of lateralized deficits¹⁶. Furthermore, “spatial” and “non-spatial” deficits in neglect have been associated with different cortical lesions, which have furthered our understanding of attentional networks in the brain. To summarize, HN is a stroke syndrome that causes hemi-inattention which effects both spatial (or lateralized) and non-spatial aspects of attention.

2.1.2 The Neuroanatomy of (In)Attention:

Typically, HN patients sustain damage to their right hemisphere and neglect the left hemifield. Cortical and subcortical lesions at many locations can cause HN. These include the

inferior parietal lobule, superior temporal gyrus, middle frontal gyrus, inferior frontal gyrus, arcuate fasciculus and superior longitudinal fasciculus²⁸⁻³⁰. Animal studies have implicated subcortical structures like the hypothalamus³¹. fMRI studies of HN patients reveal disrupted dorsal and ventral attention networks (DAN and VAN)²⁴. Compared to healthy subjects, Inter-hemispheric functional connectivity in these networks decreased in HN patients, while intra-hemispheric activity between networks increased^{32,33}. HN lesions often coexist with visual, motor, limbic and frontal lesions, which complicates treatment and diagnosis.

The dorsal attention network (DAN) includes the Frontal Eye Fields (FEF), sensorimotor cortices and parietal reach regions. It is involved in preparing and executing top-down, or goal-oriented, attentional shifts. The ventral attention network (VAN) includes the Fusiform Face Area, the amygdala and other fronto-ventro-temporal structures. This system is specialized for detection of behaviorally salient stimuli and is lateralized to the right brain^{9,11}. Lesions that cause HN usually affect both dorsal and ventral streams functionally, but damage to one system may interact with the other^{23,34}.

Direct evidence for the locus of spatial attention comes from a neurosurgical experiment³⁵. During tumor resection surgery, the superior occipitofrontal fasciculus (SOF) was stimulated during a line bisection task. In this task, a subject is asked to divide a horizontal line in half with an orthogonal line. Biases in spatial awareness are measured by how far away from the midline the subject's bisection lies. In this experiment, the subjects' bisections were shifted to the right when the SOF was stimulated. This suggests that neural information transferred between frontal and parietal regions is integral for spatial awareness or attention. This finding references well with a third resting state network, the fronto-parietal network (FPN), which is implicated in goal-directed cognition³⁶.

To summarize, HN can result from lesions to the DAN, VAN and FPN networks in the brain. These networks contribute to localization of attention, identification of stimuli and goal-directed behaviors respectively. Together, these begin to explain the constellation of deficits found in HN. Patients have difficulty orienting towards relevant stimuli, processing them and responding to them. They also have difficulty maintaining attention to complete tasks. Unlike the motor network, attention spans multiple lobes and networks in the brain, underscoring its distributed functionality in the brain.

2.1.3 Hemineglect Rehabilitation and a Rationale for BCI

Before developing control features for an HN BCI, we first reviewed current rehabilitation strategies and provide a rationale for why a BCI approach will work. Rehabilitation strategies for HN have been divided into top down and bottom up approaches. Top down approaches develop strategies to compensate for deficits. For example, repeatedly instructing a patient to remain alert improves HN deficits²⁴. Bottom up approaches improve symptoms by altering the perceptual experience of the patient. These include using prism goggles, pouring cold water in the ear and virtual reality approaches (see Appendix A for a review of rehabilitation approaches). A BCI approach to HN rehabilitation is likely to succeed for five reasons. First, existing interventions demonstrate how attentional deficits are transiently recoverable^{7,14,37}. Second, BCI approaches make use of computer games that can combine top-down and bottom-up methods. Third, many patients can eventually recover from neglect⁷, which again suggests that the deficit is recoverable. Fourth, BCI approaches can be administered at home and dosed more regularly than interventions that require a therapist. Fifth, successful BCI treatments to improve attentional deficit in attention-deficit hyperactivity disorder exist³⁸, which suggests other attention deficits are recoverable with similar methods. The first step in this BCI

approach is to create a model of how electrophysiological signals represent attention. This itself is a challenge because its relatively difficult to visualize attention. To help with this, we use the Posner task, which measures both spatial and non-spatial aspects of attention. In Chapter 4, we will use the Posner task to develop control signals for BCI rehabilitation.

2.1.4 Measuring Attention: The Posner Task

Michael Posner begins his work “Orienting of Attention” struck by “the idea that a hidden psychological process like the formation of a thought might be rendered sufficiently concrete to be measured”²⁶. Measuring hidden psychological processes is perhaps a prerequisite to Mitchell’s vision, quoted at the beginning of Chapter 1. Posner’s goal was to use the concept of spatial attention to corroborate human psychological experiments with physiological animal experiments. He broke down the process of responding to cued stimuli into four steps.

Orienting is the aligning of attention with the source of sensory input or internal semantic structure stored in memory

Detecting is when a stimulus reaches a level of representation in the nervous system where the individual can report its presence

Locus of Control (Intrinsic/Extrinsic) is whether orienting was caused by an external stimulus (e.g. a loud sound behind you) or out of one’s own volition (e.g. purposefully reading this manuscript)

Covert attention is attending to a location in space where your eyes are not fixated (i.e. looking out of the corner of your eye)

Posner used these concepts to analyze a cued spatial attention task. In the task, a participant began every trial by fixating on a central location. A central cue appeared that indicated whether

a stimuli would appear on the left or the right of screen. Most of the time, the cue indicated where the target would appear (valid trials). This ensured that the participant attended to the cued location. On a small number of trials, however, the cue pointed one direction and the target appeared in the opposite location (invalid trials). Posner found, that valid trials yielded faster reaction times than invalid trials. Posner found that he could measure where attention was being allocated by quantifying the time necessary to reallocate it. When keeping eyes centrally fixated, the participant shifted her “spotlight of attention”, independently from the location of eye fixation, toward the cued location. On the rare invalid trial, when the target appeared in an uncued location, the participant was forced to re-orient attention to the uncued location. Longer reaction times quantified this unseen cognitive maneuver. This experiment validated the idea that attention is limited: when attending to a location, one is simultaneously *not* attending to another location. No surprisingly, the deficits experienced by HN subjects can also be measured with the Posner task.

Rengachary et al. found that the Posner task captured both spatial and non-spatial aspects of HN. First, HN subjects were slower to respond to trials in their neglected visual field. Healthy subjects respond with equal RTs to targets on either side of their visual field. Second, HN subjects are slower to respond to all stimuli compared to healthy controls, regardless of which side it was on. Furthermore, these differences scaled with chronicity of the injury. Acute patients showed larger deficits in the Posner task than chronic patients³⁷. The difference between reaction times in the left and right visual field marks the lateralized, spatial deficit in HN. The increased reaction times (RTs) for both sides indexed the non-spatial deficit in HN. Furthermore, Rengachary et al. also found that the Posner task was more sensitive to detecting clinically relevant aspects of HN than standard clinical testing³⁹. In summary, the Posner task measures

attention and has been shown to measure the extent of HN deficits. In Chapter 4 we use the Posner task to interrogate the electrophysiological correlates of attention. We first review what has been previously demonstrated in the electrophysiological attention literature.

2.1.5 The Electrophysiology of Attention

Electrophysiological correlates of spatial attention have been investigated in multiple recording modalities. Single unit (i.e. neuron) recordings measure firing rates of neurons from electrodes implanted directly into the cortex. Single unit studies provide fine grain detail on how neurons change their firing rates based on changing parameters. EEG experiments study a larger network of neurons as electrodes are placed on the scalp. These experiments measure slower oscillatory activity from larger swaths of the brain than single neuron experiments. Additionally, the visual system is organized contralaterally and in topographical maps. Visual information from the left visual field is primarily decoded in the right cerebral hemisphere and vice versa. Within each hemisphere, there is a map of visual space. Experimenters use these maps to pinpoint what region of space a neuron corresponds to, then probe that region in space with stimuli in a task.

Correlates of Lateralized Attention: Bisley and colleagues found that neuronal firing in homologous (i.e. on both sides of the brain) areas of the lateral intraparietal sulcus (LIP) correlated with lateralization of covert attention⁴⁰. The LIP has been associated with tracking motion and moving eye gaze to relevant locations in space. Using a cued attention task similar to the Posner task, Bisley et al. found that when monkeys directed attention to the left, neurons in the right LIP fired at a higher rate than those in the left LIP. Thus the locus of attention correlated with the difference between left and right LIP neuronal firing rates. Similarly, EEG studies in healthy subjects have shown differential encoding of lateralized attention using alpha

power. In cued spatial attention tasks, the normalized difference in alpha activity over homologous parieto-occipital regions correlated with the locus of attention^{19,41}. The authors hypothesized this “alpha index” increases the signal-to-noise ratio of the neural control of spatial attention⁴². This signal, however, provided relatively low predictive power about the locus of attention on single-trials. Unfortunately, ECoG doesn’t cover both sides of the brain because ECoG grids are typically applied to one cerebral hemisphere. Nevertheless, these experiments provide candidate control signals for the neural correlates of lateralized attention for our studies. In summary, multimodal experimental evidence suggests that lateralization of spatial attention is encoded in differences of local neuronal activity and alpha oscillations across cerebral hemispheres.

Correlates of Reaction Time: Electrophysiological research on non-spatial aspects of attention (i.e. RT) are much broader than efforts to uncover lateralized attention phenomena. This is because reaction time (RT) measures a long sequence of neural processes. For example, a subject might take longer to identify a stimulus or that same subject might execute a slower motor response. Both will result in longer reaction times. Single unit studies have shown that the neural substrates of RT begin with single neurons in the primary sensory regions. When a monkey passively listens to auditory stimuli, the neurons in that monkey’s primary sensory regions are less active than when a monkey actively listens to the same stimuli for the purpose of responding to them²⁰. The finding that attention increases the basal firing rates of sensory neurons has been shown across sensory domains⁴³. This provides evidence that early processes in the sequence of neural events leading to a behavioral response are modulated by attention.

Historic EEG studies investigated neural activity just before the initiation of a response and discovered what’s known as the “bereitschaftspotential”. The bereitschaftspotential is alpha (8-

13 Hz) oscillatory activity that increases in power just prior to movement. This rhythm has been exploited for motor BCIs because it will remain intact even if the spinal cord is severed. If an individual is paralyzed but has an intact brain, the Bereitschaftspotential will predict when that individual desires to move. More generally, EEG correlates of reaction time have been found in the alpha and beta (13-30 Hz) frequency ranges. In a dual modality, auditory and visual, reaction time task, Senkowski et al. found that beta power correlated with RT in multiple areas in the brain, including frontal, occipital and sensorimotor cortices. Importantly, the correlation between beta and RT was negative, which means the higher the beta activity the faster the reaction time. The authors suggest that beta activity marks increased activation associated with multisensory processing⁴⁴. To summarize, RT measures a host of processes that transform the perception of a stimulus to an action. However, multiple lines of evidence suggest that the measured brain activity correlates with future reaction times.

Relationship between neural oscillations and attention: We previously discussed how alpha oscillatory activity correlates with the direction of cued attention and beta oscillatory activity correlates with RTs. Now we discuss *why* attention might be related to oscillatory activity in the first place. To do this we first discuss two additional experiments and a theory. In the first experiment, visual and audio oddball tasks were interleaved, and a monkey was required to attend only to visual or auditory stimuli. The goal of the task was to respond to a low frequency audio or visual “oddball” amongst high frequency stimuli (i.e. (auditory) respond to the rare “beep” and ignore the frequent “boop” sounds *or* (visual) respond to the rare green square while ignoring the frequent red squares). When the monkey attended to the visual stimuli, neural oscillations entrained to the timing of the task creating high delta (1-3 Hz) power and gamma (75-100 Hz) activity increased during specific phases of the delta oscillation. Gamma

activity is believed to represent local neuronal firing. This effect only occurred in the visual cortex when the monkey attended visual stimuli. When the monkey attended auditory stimuli, the effect occurred in the auditory cortex. When attention was deployed to a sensory system, the corresponding brain region oscillated. Then, the local neuronal activity in that region became entrained to the ongoing oscillation. The authors suggest that low frequency oscillations are responsible for transiently increasing the excitability of neurons in primary sensory cortices in order to maximize perception of rhythmic stimuli⁴⁵. Delta essentially amplified rhythmic stimuli by entraining the brain to their timing.

The finding that neural oscillations relate to attention is consistent with findings that the phase of low frequency oscillations predict visual perception^{46,47}. Perception can be measured by asking subjects whether they have seen a specific stimulus or not. The phase of an oscillation is a description of where that oscillation is in its cycle. The phase starts at 0 and completes a full cycle at 360 (i.e. 2π). In multiple experiments, authors have found that the phase of ongoing theta (3-7 Hz) or alpha oscillations predicts whether a stimulus is detected or not^{46,47}. If a visual stimulus is presented at the 180-degree phase of the oscillation, the stimulus is more likely to be detected. Beyond perception, interactions between low frequency phase and high frequency amplitude have also been shown to predict reaction time. One study showed that the phase locking value (PLV) between delta/theta and gamma bands correlated with reaction times⁴⁸. These findings suggest that perception and performance may depend on parcellated periods of time that are created by oscillations.

Our subjective experience of the world is a continuous one. We do not see time moving in steps or space represented as pixels. However, this is an illusion created by the brain. The visual system is represented by individual neurons in the retina that correspond to discrete sections of

the visual field. Despite this, we visualize a continuous visual field. Why then, should time be any different? Discrete attentional sampling was proposed in the “Active Sensing” hypothesis¹³. The authors suggest that humans may perceive our environment similarly to robots that discretely sample signals from the environment. The strongest example of this is in olfaction where sensing a smell is intimately linked with respiration. When a mouse inspires, it is more likely to smell something salient, therefore the sensation of smell is not passive, but actively generated. Visual attention can be moved independently of eye muscles, making the active sensing hypothesis less obvious in visual attention. However, the findings that perception depends on the phase of ongoing oscillations suggests that cognitive rhythms may underlie visual sampling. Critics of this hypothesis suggest that the $1/f$, or scale-free, distribution of brain activity allows for continuous temporal sampling⁴⁹. These theoretical arguments underscore the difficulty in understanding subjective human experience. However, “experience” is one way to define consciousness and consciousness is a cognitive state that we can manipulate and measure.

2.2 Consciousness

2.2.1 Defining Consciousness

There are at least two ways to define consciousness, “experience/awareness” or “arousal/wakefulness”, and both relate to attention. In hemineglect, both the awareness of space and general levels of arousal are affected. In Chapter 5 we will report novel findings on the electrophysiological correlates of slow wave sleep and Propofol-induced loss of consciousness. Even though arousal and experience are both reduced in unconscious states, we will review these terms separately at first to better understand the meaning of consciousness.

Awareness: Many have had the experience of driving while thinking about something other than driving. If a driver's attention is directed to sending a text message or singing a song, he will be less aware of his environment. However, a driver can lack awareness even while looking at the road, provided something else occupies his thought. Awareness references the subjective experience of reality, and is difficult to measure. For example, patients with "locked-in syndrome" (LIS) have functional brain activity, but are unable to move their body. LIS is not a disorder of consciousness, it is a disorder of the motor system. However, it might be difficult for someone to tell the difference between an LIS patient and a patient without brain activity. An evolving clinical understanding of "awareness" comes from differentiated states like LIS, minimally conscious states or vegetative states, typically after a brain injury⁵⁰.

The criteria for a coma or a vegetative state includes no evidence of awareness of self or environment, an inability to interact with others and no purposeful or voluntary responses to stimuli. Unlike the participants in the tasks described in previous chapters, a patient in a vegetative state cannot respond to stimuli. However, vegetative patients awaken from sleep, and their bodies can survive with assistance. In contrast, patients in a minimally conscious state (MCS) have unequivocal evidence of awareness. These patients respond to stimuli, follow simple commands and are aware of their environment. However, these patients may not resemble their uninjured selves. They are limited in their cognitive capacity, which leads to deficits in communication, sustained attention and abstract thought, to name a few. Here we see that awareness is defined by the ability to respond purposefully to stimuli. This definition relates to our attention tasks where we measure purposeful responses to stimuli. Unfortunately, while MCS patients are undoubtedly aware, they are minimally aware and suffer deficits in most measurable

cognitive functions. Interestingly, studies have shown that zolpidem (aka ‘ambien’, a GABA agonist like Propofol) can transiently recover lost cognitive functions in MCS patients⁵⁰.

Arousal: Another common experience in humans is waking up from sleep. This behavior is linked to our homeostatic regulation governed by the autonomic nervous system. The autonomic nervous system responds to danger and is key for survival. When we awaken our digestive system activates, our heart rate increases, hormones are released and we are generally readier to address our environments. Clinically, coma states are considered the lowest arousal states. Coma patients won’t open their eyes or respond to unpleasant stimuli. However, when a patient awakens from a coma, they either open their eyes or exhibit brainstem reflexes. Unlike in coma, patients in vegetative states exhibit spontaneous eye opening and reflexes, suggesting they are awake, without eliciting signs of awareness. However, in most states of consciousness that healthy individuals are familiar with, including sleep, arousal and awareness scale together.

2.2.2 Sleep

Many neurophysiological events occur during sleep that affect our waking behaviors. For example, memories are consolidated during sleep and sleep restores our capacity for attention and awareness⁵¹. Furthermore, sleep may help our brain clear waste material generated by neurons during the day⁵². Some researchers believe that sleep developed alongside the ability to learn and pay attention. This theory is supported by phylogenetic evidence. Animals like *Caenorhabditis elegans* (*C. elegans*) cannot engage in operant conditioning (i.e. learning from trial and error) and only sleep prior to developmental events like molting. Fruit flies, on the other hand, sleep and learn more similarly to humans. For example, if you prevent a young fly from sleeping, it develops lasting cognitive deficits. Thus, sleep may be a counterweight to higher cognitive functions like attention⁵³.

Sleep also shares some similar qualities to attention. When we sleep, information flow across our cortex becomes limited⁵⁴. This is not unlike what happens in selective attention experiments discussed previously. For example, in the dual auditory-visual oddball task discussed previously, when a subject attends to an auditory stimulus, but not a visual stimulus, the visual attention system is quiescent⁴⁵. Similarly, different parts of the brain can shut down selectively during sleep⁵⁵. What's more, sleep is a waxing and waning process with multiple stages that have different characteristics. Certain neural connections can increase during sleep as well, such as the functional connections between the cortex and the hippocampus that promote memory consolidation⁵⁶. It may be energetically advantageous to shut down some neural connections to promote others. Historically, sleep stages have been defined by EEG activity and eye movements.

Slow wave sleep (SWS) is differentiated from rapid eye movement (REM) sleep based on these EEG and eye movement definitions. Dreaming is thought to occur during REM sleep, therefore the brain is relatively active. SWS is a more quiescent state and is characterized by large delta oscillations in EEG. These oscillations correspond to "UP" and "DOWN" states. During UP states, cortical neurons are more likely to fire than during DOWN states⁵⁷. In this manner, SWS regulates neuronal activity across time. During DOWN states, there is less activity in neurons which might allow for important sleep functions like cellular repair. In contrast, UP states may facilitate memory consolidation. Interestingly, this is not unlike the interaction between brain regions in the dual oddball task where attentive UP states occurred during behaviorally relevant time periods. Another potential reason for intermittent UP states during sleep is so that environmental awareness is maintained. Sleeping humans are arousable during sleep and can even respond to stimuli⁵⁸, presumably so they can avoid danger. During general

anesthesia, however, humans are not arousable. Sleep shares similarities and differences from induced unconscious states during anesthesia, in Chapter 5 we will focus on comparing SWS to Propofol-induced loss of consciousness.

2.2.3 Propofol Induced Loss of Consciousness (PILOC)

Propofol is a gamma-Aminobutyric acid (GABA) agonist, that is commonly used to cause sleep and amnesia during surgery. It was also the cause of Michael Jackson's untimely death. Propofol has several qualities that appear similar to sleep. For example, when someone hasn't slept for more than 24 hours, sleep debt accrues. Sleep debt can be relieved by PILOC. Furthermore, the greater the sleep debt accrued, the faster Propofol takes effect^{59,60}. However, unlike sleep, patients undergoing PILOC are not arousable. In this sense PILOC is a deeper state of unconsciousness than sleep. The neurochemistry of GABA helps explain Propofol's mechanism of action.

GABA is the main inhibitory neurotransmitter in the brain. The neurons that produce GABA in humans typically have inhibitory actions. This means that they prevent other neurons from firing by changing their cellular membrane potentials. GABA receptors are in many places in the brain, however, studies suggest that the main sites of action for Propofol are regions controlling arousal, associative functions and autonomic control⁶¹. This is likely promoted by different receptor concentrations and accessibility of the drug through the vasculature.

EEG studies have shown similarities between PILOC and SWS as both unconscious states have slow delta waves⁶². These slow waves are believed by some to serve as a pacemaker for the UP and DOWN states⁶³. However, these slow waves appear more disorganized in PILOC than in SWS. A major difference between SWS and PILOC is the occurrence of alpha power

frontalization that only occurs in the latter⁶⁴. Although sleep has alpha-like activity in the form of sleep spindles, alpha frontalization is unique to GABA-related loss of consciousness and presumably caused by differences in activity due to inhibition of the frontal cortex and thalamus, which are reciprocally connected. As previously mentioned, GABA agonists, such as Zolpidem (aka Ambien), which is similar to Propofol in its mechanism, have been shown to awaken people from minimally conscious states⁵⁰. Zolpidem is a GABA agonist that is used to aide sleep.

3) Methods: Measuring Phase-Amplitude Coupling

3.1 From Neuronal Firing to Scalp Recordings

Neurons are cells that maintain a difference in charge stored on their cell membranes, a polarization. They fire “action potentials” or “spikes” by rapidly depolarizing their membrane potentials relative to the extracellular environments. ECoG (and EEG) measures the depolarization patterns of groups of neurons. Groups of neurons from one brain region can spike in concert, or spike in a disjointed fashion ⁶⁵. When they spike together, their electrical waveforms are summed to make a larger waveform that is detectable on ECoG. When they fire disjointly their independent waveforms cancel out.

To better understand this process, imagine a packed baseball stadium where fans are analogous to neurons. If you stand in a building across the street from the stadium, but you had a microphone on a long enough boom arm, you could use it to listen to one individual talking in the stadium. This is similar to recording spikes from single neurons in the brain. Without the long boom arm, you can still record stadium sounds from your distant position. When the home-team hits a home run, and the whole stadium erupts in jubilation, the aggregate sound from fans cheering in unison will be recorded outside of the stadium. This is similar to an ECoG recording. The aggregate noise recorded outside the stadium is the summation of many individual voices, just like ECoG recordings are generated by the summation of neuronal activity. In this sense, global activity is generated by local activity. However, if we continue with this analogy we’ll find that the opposite is also true. Local activity can be recruited into a global trend. For example, a couple that is deep in conversation prior to a home run will be jolted out of that

conversation by the home-run cheering crowd. They may even forget their conversation and join the cheer. Similarly, the activity of an individual neuron can be modulated by the activity of surrounding neurons through their effect on the extracellular environment. In this analogy, oscillatory activity is analogous to the organized “wave” that commonly occurs during athletic events². These are typically started by a passionate group of individuals, but successful waves recruit many more participants. Interestingly, the EEG analog was seen in the very first recordings of brain activity.

Why we study neural oscillations: Describing the brain in terms of oscillatory activity originated from the first EEG recorded by Hans Berger in 1929. Berger was the first to use galvanometers to record brain activity (which he believed was psychic energy) from healthy and brain damaged patients. Notably, he devised experimental paradigms that still exist today in renewed form. For example, when recording EEG from a paralyzed patient injured by a gunshot wound to the brain, he unexpectedly fired a pistol behind the subject to measure changes in brain activity caused by “involuntary attention”⁶⁶. This is like the invalid trials in the Posner task, which cause involuntary shifts in attention, albeit in a far less cruel manner. In his early recordings Berger noticed waves with roughly 120-180 millisecond spacing and waves with roughly 30-45 millisecond spacing. These became known as alpha (~10 Hz) and beta (~20 Hz) waves⁶⁷. To this day it is not entirely known why oscillations exist in the brain. However, most biological processes, from cell division to breathing, are cyclical or oscillatory. One theory suggests that neuronal oscillations are a result of the brain’s need to conserve energy and oscillations are an energetically efficient way of creating complex brain interactions⁴⁹. Recent

² According to Wikipedia, “the wave is an example of metachondral rhythm achieved in a packed stadium when successive groups of spectators briefly stand, yell and raise their arms.”

publications have suggested that oscillations in the brain may not be as sinusoidal as traditionally believed^{68,69}. In this work we will be careful to distinguish between oscillations in filtered brain recordings and non-sinusoidal activity that occurs in raw signals recording brain activity. The difference between these two lies in the methods used to measure oscillations.

Quantifying oscillations with Wavelets: Measuring oscillations in the brain typically takes a signal in the “time” domain and transforms it into the “frequency” domain. To do this we define a time period for analysis, which limits the possible frequencies that can be measured. For example, if we look at a 1 second period of data, the lowest frequency we can detect is a 1 Hz oscillation. Similarly, if we sample that 1 second period 1000 times, then the fastest frequency we can detect is a 500 Hz signal. Furthermore, most frequency decompositions fit sinusoids to the data even if the data are not sinusoidal. These are mathematical principals that apply to frequency-domain analyses. There is a tradeoff between time resolution and frequency resolution. To measure low frequencies, we require long intervals of time. But the longer the period we analyze, the less we can say about when an oscillation occurs. The time-frequency tradeoff says that the more precision we have in measuring a neural event in time, the less precision we have in measuring its frequency content. We acknowledge this limitation and use a wavelet approach to balance the time-frequency tradeoff. Wavelets are wave-like oscillations that have both a temporal beginning and end, as well as a sinusoidal component at a chosen frequency. The formula for the Morlet or Gabor wavelet is as follows:

$$\Psi_{f_0, \sigma}(t) = \frac{1}{2\pi\sigma} e^{\frac{-t^2}{2\sigma^2}} * e^{i(2\pi f_0 t)}$$

The first factor in the equation is a Gaussian function with a standard deviation (σ). When the Gaussian function is transformed from the time-domain to the frequency-domain it has

the special property of remaining a Gaussian function. The second factor is Euler's identity with a frequency f_o .

$$e^{ix} = \cos x + i \sin x$$

When this wavelet is convolved with an ECoG signal, the output is another signal that is the same length as the original signal. The output, or wavelet decomposition, will have higher values when the original signal has high amplitude in a f_o frequency range (Figure 3.1.1).

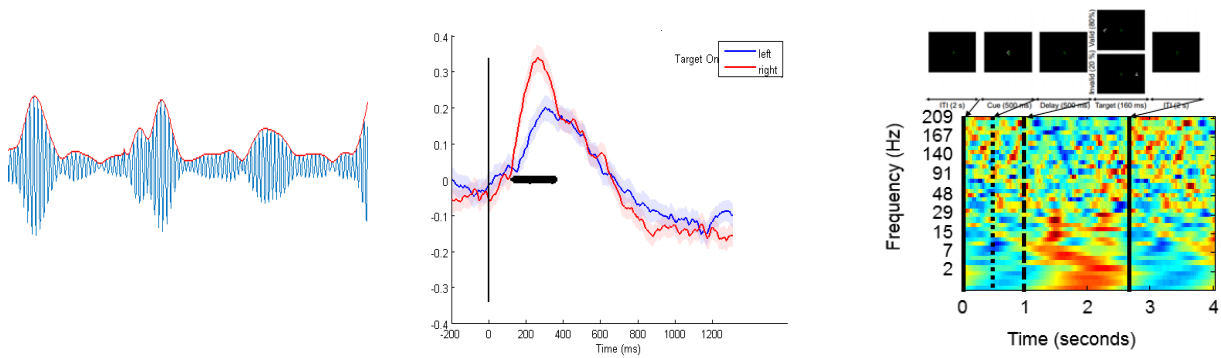


Figure 3.1.1: Signal processing and analysis methods: (Left) Output of a wavelet decomposition in the gamma (75-100 Hz, blue) with amplitude envelope (red). Increased firing of single neurons influences the increases in gamma power. (Middle) Averaged gamma amplitude for left and right targets shows how right targets have higher gamma power. (Right) A spectrogram of 1-209 Hz showing all frequencies where red means higher power and blue means lower power.

The width of the wavelet, which is equivalent to the standard deviation (σ) of the Gaussian functions, determines how many cycles of the f_o frequency the wavelet contains. Here the time-frequency tradeoff emerges again. The wider the wavelet is in the time domain (i.e. the larger the σ), the more accurate it is at detecting the f_o frequency. However, larger σ also means

the wavelet decomposition will be less temporally accurate. This property becomes important to the topic of phase-amplitude coupling, which we discuss in the next chapter. First, we discuss phase and amplitude.

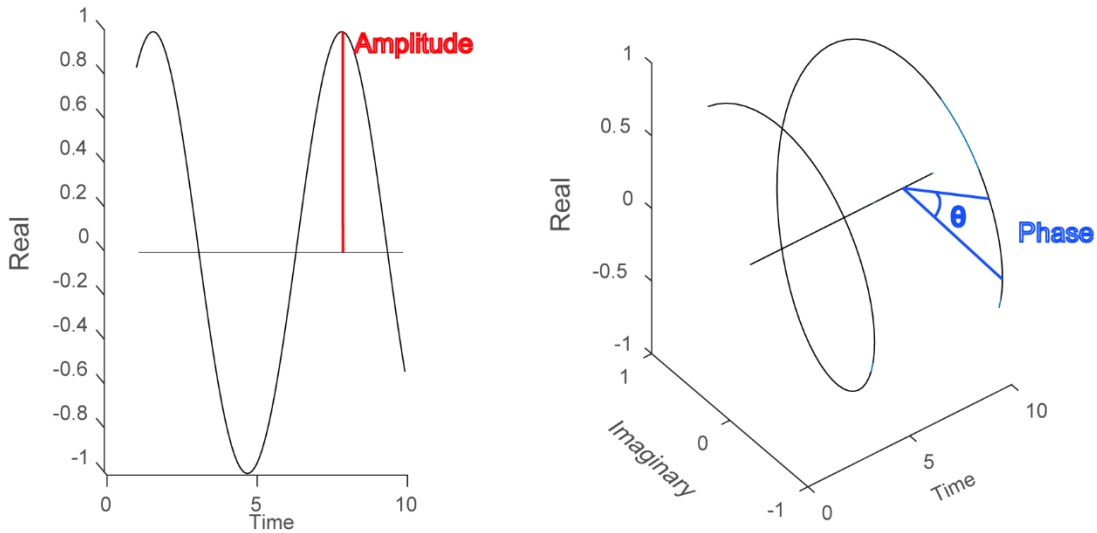


Figure 3.1.2: Amplitude and phase of from a complex signal. (Left) A representation of an oscillation in the real and time domains demonstrates amplitude (red). However, multiple locations on the oscillation have the same amplitude. (Right) Phase is the angle of the wave in the imaginary and real domains (blue). Phase and amplitude together allow us to precisely locate any point in the oscillation.

Euler's formula provides the connection between mathematical analysis and trigonometric functions representing oscillations. While we commonly discuss the amplitude of oscillations, like the volume of our music, we less often discuss phase (**Figure 3.1.2 right**). Phase is useful because it tells us how far an oscillation is along its cycle, independently of its frequency. Its values range from 0 to 2π , completing a full circle or cycle. In phase-amplitude coupling, the amplitude envelope of one frequency couples to the phase of another frequency.

3.2 Phase-amplitude coupling (PAC)

Phase amplitude coupling (PAC) is when the amplitude of one frequency band preferentially increases on particular phases of a second frequency band. It is analogous to amplitude modulation (AM) radio. When a listener tunes into the AM radio station 1010 WINS, she tunes into a 1010 KHz carrier frequency. If a musician on the station plays the note A at 440 Hz (i.e. the modulating frequency), then the amplitude of the 1010 KHz carrier frequency is modulated at 440 Hz. This means that at a particular phase (e.g. 0 phase) of the modulating frequency (440 Hz), the carrier frequency (1010 KHz) amplitude will be maximal. At the opposite end of the modulating frequency's cycle (e.g. 180-degree phase), the carrier frequency amplitude will be minimal. Your speakers play the amplitude envelope of the 1010 KHz signal, which will ultimately sound like the note A at 440 Hz. A similar process has been found in neural signals and it is called PAC.

In neural signals, the purpose of PAC is less clear than its radio equivalent. Early studies found that EEG patterns in mice were linked closely to breathing patterns⁷⁰. More behaviorally relevant olfactory information is available during inspiration, compared to expiration. Many believe that neural PAC differs from its AM radio analog because the information is carried in the high-frequency content (i.e. neuronal firing), rather than the low frequency content (i.e. breathing cycle). In AM radio, the information is the low frequency content (i.e. the 440Hz modulating frequency). PAC in the brain has been correlated with attention¹³, learning⁷¹, and memory⁶⁹. Recently, PAC has even been shown between the brain and gut oscillations⁷². Since most biological processes, from cell division to neuronal firing, are cyclical, it is not surprising

that some biological cycles influence other biological cycles at different time scales. In the brain, however, it is not yet clear if PAC affects cognition and behavior, or if it is an epiphenomenon of some other process.

3.3 Measuring PAC

There are many ways to measure PAC, but we chose to use the modulation index (MI) because it accurately measures the intensity of coupling and performs better than other measurements. For a review of different methods and their strengths see Tort. et al. “Measuring phase amplitude coupling in Neural Oscillations of Different Frequencies”⁷³. The first step in calculating MI is constructing a phase-amplitude plot. To calculate the phase-amplitude coupling between 8 Hz phase and 28 Hz amplitude, we first divide the 8 Hz signal into bins (i.e. 20 bins from -180 to 160, 160 to 140, etc.). Next, we collect all the amplitude envelope values of the 28 Hz signal that correspond to the phase bins of the 8 Hz signal. In **Figure 3.2.1** we see the amplitude signal (i.e. 28 Hz) in plot **B**. The phase of the phase signal (i.e. 8 Hz) is in **Figure 3.2.1, D**, colored from $-\pi$ to π with rainbow colors. We project the colored phase bins of the 8 Hz signal in **Figure 3.2.1, D** onto the amplitude envelope of the 28 Hz (**Figure 3.2.1, B**). We collect all of the amplitude frequency samples that correspond to a phase bin (or color) and average them. We plot these averages by phase bin to make a phase-amplitude plot (**Figure 3.2.1, C**). When we normalize the phase-amplitude plot by the sum of all amplitudes it becomes a probability density where the normalized amplitudes sum to 1. If there was no coupling between the 8 Hz and 28 Hz frequencies then the phase plot would look like the uniform distribution (gray line). However, we see that the observed distribution differs from the uniform distribution (**Figure 3.2.1, C, red dots vs grey dashed line**), which means coupling exists. To measure the difference between the observed and uniform distance we use the Kullback-Leibler

divergence (See Chapter 4, methods). This quantifies the intensity of coupling. We can repeat this procedure for every pair of frequencies under investigation to generate a *comodulogram* (Figure X, E). The comodulogram identifies which frequency pairs are coupled and how intense the coupling is between them.

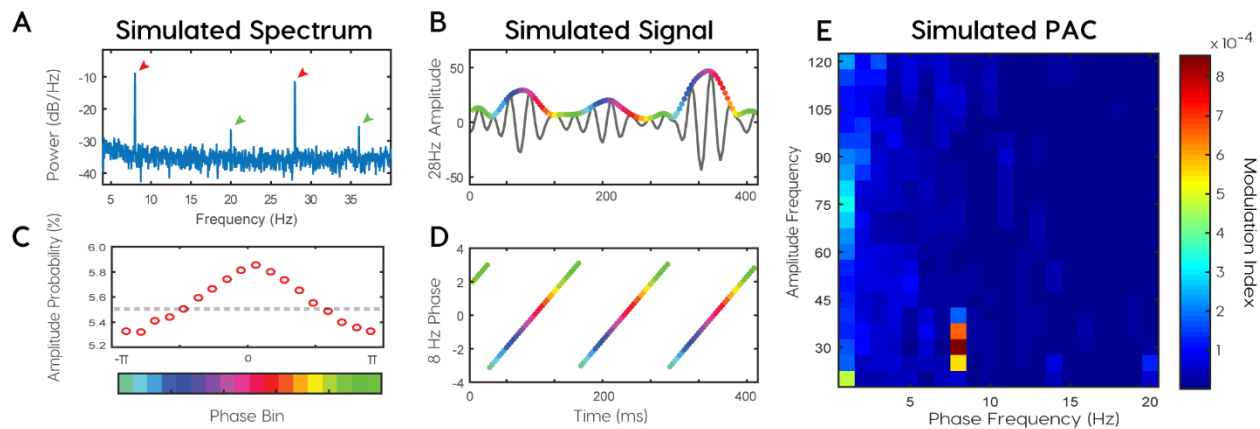


Figure 3.2.1: Phase amplitude coupling measurement with modulation index.

One important consequence of coupling is the existence (or creation) of two new frequencies, the sum and the difference of the frequencies that are coupled. If an 8 Hz phase is coupled with a 28 Hz amplitude then two new frequencies must exist at 20 Hz and 36 Hz (Figure 3.2.1, A). Previously we mentioned how the standard deviation of the wavelet determined its spectral precision. To capture coupling phenomena, too much precision is a bad thing. This is because the wavelet that decomposes the amplitude signal (i.e. the 28 Hz signal) must also include the additional frequencies (20 Hz and 36 Hz). If it does not include these “side peaks” then no coupling can be measured between 8 Hz phase and 28 Hz amplitude⁷⁴. This means that the bandwidth of the amplitude wavelet must be twice the frequency of the phase wavelet.

Therefore, the bandwidth required to see 8 Hz-phase coupling is 16 Hz and our amplitude wavelet for 28 Hz-amplitude, must include 20 Hz to 36 Hz. This relationship led us to design custom wavelets that improved our resolution of PAC phenomena with high-frequency (>4 Hz) phase. Custom wavelets were the most likely reason we found PAC phenomena that were unreported in comparable attention tasks^{48,75}. Separate wavelet libraries are justified because spectral precision is important in low frequency phase signals, but spectral imprecision is required in high frequency amplitude signals. **Figure 3.3.1** shows the differences in resolution when we widen high frequency amplitude bandwidths from 3 Hz to 20 Hz (**Figure 3.3.1, left vs right**)

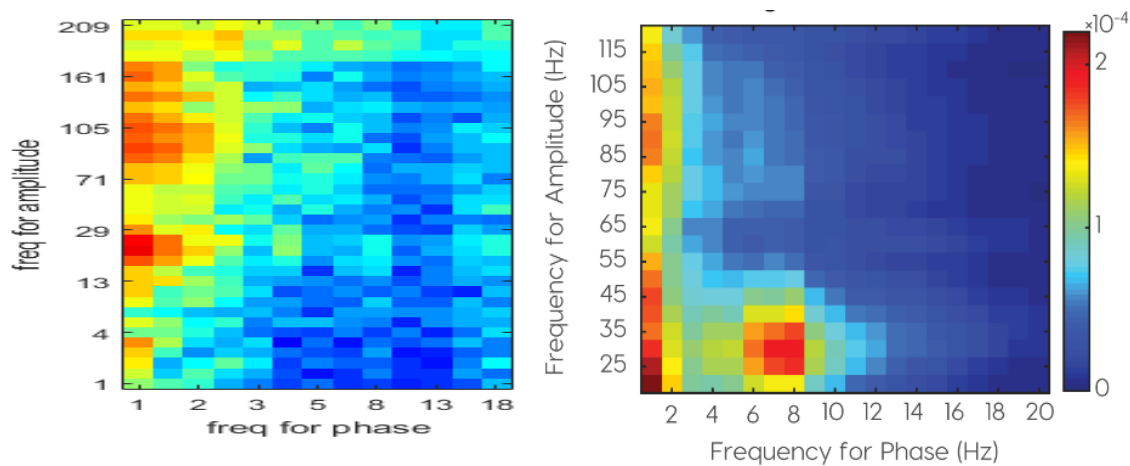


Figure 3.3.1 Importance of high frequency for amplitude signal wavelet bandwidths. (Left)

A comodulogram with 3 Hz wavelet bandwidths for amplitude and phase signals. (Right)

Comodulogram with 0.8 Hz phase signal bandwidth and 20 Hz amplitude signal

bandwidths. The wider amplitude signal bandwidths allow for resolution of higher phase

PAC phenomena at 6-8 Hz phase

3.4 Theory of PAC in neural oscillations

Two hypotheses are relevant to PAC and cued spatial attention (for an extensive review see (Bonnefond, Kastner, and Jensen 2017)). The communication through coherence (CTC) hypothesis proposes that excitability windows of two communicating brain regions are temporally aligned by low frequency oscillations to promote information transfer across regions (Bastos, Vezoli, and Fries 2015; Fries 2005). To understand how the CTC hypothesis requires PAC to impact inter-regional communication, consider two communicating regions of the brain A and B. If these two regions both have strong 1 Hz oscillations, then local neuronal excitability also oscillates at 1 Hz. A neuron in region A is more likely to fire during an excitable phase in region A. It is also more likely to excite a neuron in region B because signals from region A reach region B during an excitable phase in region B. In this model PAC must exist in both region A and B because temporal windows created by low-frequency oscillations will be filled with the transmitted signal. In contrast, the gating by inhibition (GBI) hypothesis proposes that low frequencies periodically suppress information in a neuronal population (Jensen and Mazaheri 2010). The larger the power of the low frequency oscillation, the smaller the temporal window of excitability. This hypothesis suggests oscillations locally inhibit neural activity and no oscillatory information from region B is required to understand if region A is communicative. There have been efforts to unify these hypotheses under a common theoretical framework (Bonnefond, Kastner, and Jensen 2017), but questions persist about the mechanisms that relate PAC to cognition. In our study we find evidence for a suppressive role for PAC in the theta-alpha-phase/beta-low-gamma amplitude range, which fits with the GBI hypothesis (Chapter 4).

3.5 Challenges to PAC theory and measurement: Non-sinusoidal, Sharp Waves

Several issues with phase-amplitude coupling analyses have been reported. First, most methods of measuring neural oscillations and PAC assume that the raw signal is stationary. That is, the raw signal's mean and variability don't change over time. This is not true in neuropsychological studies because experimenters purposefully perturb cognition at predetermined times to measure a changing signal. These non-stationarities may appear as power in multiple frequencies when the signal is decomposed into individual frequencies. When raw signals have non-stationary or non-sinusoidal activity it is possible to measure phase amplitude coupling between two frequencies when there is little other evidence for those two frequencies interacting. One specific type of non-sinusoidal wave recorded from brain activity is the “sharp wave”

Sharp waves are non-sinusoidal waves that look like intermittent spikes rather than oscillations. Sharp waves have been shown in epilepsy patients and in the hippocampus, but have been rarely discussed in the attention and consciousness literature. Recently Vaz et al. distinguished two forms of PAC in a memory task⁶⁹. The first was between theta phase and high-gamma amplitude, which they called “nested oscillations”. The second type of PAC was between theta-alpha-phase and low-gamma amplitude, which they called “sharp waveforms”. We emphasize this distinction because it clarifies the challenge to phase amplitude coupling analysis. A “nested oscillation” is what is traditionally believed to create PAC and many of the early PAC findings were in the delta-theta-phase coupled with high-gamma-amplitude. However, we propose that “nested oscillations” are a special case of PAC. When we look at higher-frequency-

phase and lower-frequency-amplitude (e.g. theta-beta coupling), the coupling in the raw signal looks different. It was not previously clear if sharp waves are themselves coupled or if sharp waves cause coupling across multiple frequency bands. We show that the former is true in Chapter 4 and believe that this conflict is a matter of semantics. “Sharp waves” are another special case of PAC, since they occur at regular intervals.

3.6 Evidence for Phase-amplitude coupling in Attention and Consciousness

Strong evidence for the importance of PAC in attention came from the “oddball” task discussed in the last section of 2.1.5. However, two more experiments explored PAC in attention but differed in their conclusions. The first was a cued detection task with distractors conducted by Szczepanski and colleagues⁴⁸. Subjects were cued to the left or right of a screen and told to respond as fast as possible when they saw a blue dot while ignoring distractors, which were red dots. The researchers found that PAC between delta and high gamma during the cue period correlated most strongly with reaction time when the subject was cued contralateral to the ECoG grid. This suggested that PAC was facilitating visual attention because vision is represented in the contralateral hemisphere (i.e. left stimuli are best represented in the right cerebral hemisphere). The authors also showed that event-related potentials (ERPs, a form of non-stationary brain activity) did not co-occur with PAC phenomena, suggesting that ERPs were not the cause of the PAC they witnessed.

In contrast, Esgheai and colleagues found that visual attention decreased PAC in local field potentials recorded from the Lateral Intraparietal Cortex (LIP). In a cued attention task, PAC decreased when the monkey was cued to the contralateral receptive field of the LIP being recorded. The authors concluded that high PAC meant that neurons in the LIP were more

correlated with each other. When neuronal firing is correlated, less information can be stored or processed. To illustrate this, imagine all neurons in a region oscillated together. In this case, you can predict the firing rate of one neuron with another neuron. Therefore, each neuron does not encode independent information. If, however, neurons in a region fired independently, then more information can be stored between the neurons. The same would be true for bits in a computer processing unit, if they all did the same thing less computation could get done.

There are significant differences between the experiments that could explain many of the differences. For one, local field potentials (LFP) are recorded within the cortex while ECoG is recorded at the surface. LFP recordings in monkeys are very accurately placed in the cortical region under study, unlike ECoG that is meant to record a large cortical area. Esghaie et al. took advantage of this by isolating the receptive field (i.e. the area in visual space that corresponds precisely to the brain region under study) of area LIP and using that receptive field for their stimulus presentation. This is important because ECoG records from regions surrounding the receptive field being probed. These surrounding regions can do the opposite of the region primarily associated with a receptive field. A second potential reason for differences in the findings is that the authors focused on different frequencies that could perform different functions. We disentangle frequency-specific PAC phenomena in Chapter 4.

4) Distinct Phase-Amplitude Couplings Distinguish Cognitive Processes in Human Attention

4.1 Abstract

Spatial attention is the cognitive function that coordinates the selection of visual stimuli with appropriate behavioral responses. Recent studies have reported that phase-amplitude coupling (PAC) of low and high frequencies covaries with spatial attention but differ on the direction of covariation and the frequency ranges involved. We hypothesized that distinct phase-amplitude frequency pairs have differentiable contributions during tasks that manipulate spatial attention. We investigated this hypothesis with electrocorticography (ECoG) recordings from participants who engaged in a cued spatial attention task. To understand the contribution of PAC to spatial attention we classified cortical sites by their relationship to spatial variables or behavioral performance. Local neural activity in spatial sites predicted spatial variables in the task, while behavioral sites predicted reaction time. We found two PAC frequency clusters that covaried with different aspects of the task. During a period of cued attention, delta-phase/high-gamma (DH) PAC predicted cue direction in spatial sites. In contrast, theta-alpha-phase/beta-low-gamma-amplitude (TABL) PAC robustly predicted future reaction times in behavioral sites. Furthermore, TABL PAC corresponded to behaviorally relevant, sharp waveforms that coupled to a 7.2 Hz rhythm. We conclude that TABL and DH PAC correspond to distinct mechanisms during spatial attention tasks and that non-sinusoidal sharp waves are elements of a coupled dynamical process.

4.2 Introduction

Spatial attention defines a set of cognitive mechanisms that select behaviorally relevant visual information while filtering out behaviorally irrelevant information²⁶. Attention facilitates visuomotor coordination through modulation of neural activity in visual^{17,76,77}, parietal and prefrontal regions during visuomotor tasks⁷⁸⁻⁸⁰. Phase amplitude coupling (PAC) has been proposed as a mechanism underlying attention^{48,75,81,82}. PAC quantifies the relationship between the phase of a low frequency signal and the amplitude envelope of a high frequency signal. It has been hypothesized that low-frequency oscillations serve as temporal reference frames for higher frequency (>20 Hz) activity^{45,83-85}. These hypotheses are supported by evidence that stimulus perception depends on the phase of ongoing oscillations⁴⁷. However, oscillatory hypotheses have been challenged by evidence suggesting that non-sinusoidal waveforms cause spurious PAC^{86,87}.

Two relevant PAC experiments on cued spatial attention resulted in opposing conclusions. In a human cued target-detection task with distractors, Szczepanski et al. found delta-theta-phase/high-gamma amplitude (2-5 to 100-150 Hz) PAC correlated with reaction time (RT) in cortical sites associated with the dorsal attention network. Correlations were stronger when subjects attended to the receptive hemifields of recording sites⁴⁸. The authors suggest that PAC enhances spatial attention given positive interactions between PAC, cueing direction and reaction time. In contrast, Esghaei et al. found decreased PAC between low (1-8Hz) and high (30-120 Hz) frequencies when monkeys attended to the receptive field of area MT⁷⁵. The authors conclude that PAC suppresses attention given its negative covariation with cueing direction.

Beyond differences in recording methodologies, task paradigms and species, each group focused on different low-frequency ranges (e.g. 2-5 Hz for Szczepanski et al. and 1-8 for Esghaei et al.) that may engage distinct neural circuitry. Delta (1-4 Hz) in primary sensory cortices has been shown to reflect rhythmically presented stimuli when attended⁴⁵. Theta (4-7 Hz) and alpha (8-13 Hz) have been associated with sustained attention^{88,89} and inhibition^{19,90}. Evidence suggests that delta and alpha play opposing roles in selective attention⁹¹. Finally, recent investigations into the origins of PAC revealed that distinct PAC frequency pairs correspond to waveforms that explain different cognitive processes in a memory task⁶⁹. Given these findings, we wondered whether specific PAC frequency pairs corresponded to different cognitive elements of a spatial attention task.

In this study, we investigated frequency-specific PAC phenomena during a spatial attention task and hypothesized that PAC frequency pairs would show distinct functional characteristics. Additionally, we sought evidence for non-sinusoidal waveforms causing spurious PAC^{86,87}. Using methods defined by Tort et al. to measure PAC with the modulation index, we used a non-parametric cluster-based statistical approach to find behaviorally relevant PAC frequencies^{73,92}. We functionally classified cortical sites based on their sensitivity to spatial properties of stimuli or behavioral performance. We found that theta-alpha-phase/beta-low-gamma-amplitude (TABL) PAC predicted reaction time while delta-phase/high-gamma-amplitude predicted cueing direction. Furthermore, we developed computationally inexpensive methods to detect the non-sinusoidal correlates of TABL PAC. We found that these non-sinusoidal waveforms correlate with RT, however, they also coupled to a low frequency oscillation (7.2 Hz). Our findings show that the functional characteristics of PAC depend

critically on low frequency phase and that sharp waves are elements of a coupled dynamical process.

4.3 Materials and Methods

Subjects and Data Acquisition

The study included six human participants, of both sexes, with treatment-resistant epilepsy who were undergoing invasive electrocorticography (ECoG) to detect seizure foci. None had vision or attention deficits. The data from three subjects were analyzed with different methods in a previous experiment⁹³ the remaining data were not previously analyzed. A computer monitor was placed 20 inches away from the subject's eyes. ECoG data was recorded in the subject's hospital room from platinum clinical electrodes with 2.3 mm diameter and 10 mm spacing (PMT Corporation, Chanhassen, Minnesota). The raw ECoG signals were sampled at 1200 Hz and amplified with clinical bioamplifiers (Guger Technologies, Schiedlberg, Austria). We developed custom scripts for use with the BCI2000 software platform for task presentation and data acquisition (www.bci2000.org, Schalk et al. 2004).

Experimental Design

Subjects participated in a modified Posner spatial cueing task previously described by Daitch et al.⁹³. Subjects were cued with a centrally located arrow that pointed either left or right and appeared for 500 milliseconds. After variable cue offset and an additional delay, the target appeared for 160 milliseconds. An equivalent number of left and right targets were presented in random order. A target appeared at the cued location on 80% of trials (valid) and at the un-cued location on 20% of trials (invalid). All subjects engaged in sessions where the timing between cue offset and target was fixed. Fixed trials had a cue-target interval of 500 milliseconds. Five of

the six subjects alternated between sessions with fixed and variable cue-target interval. In variable sessions, the interval between cue offset and target onset varied between 500, 1000 and 1500 milliseconds with equal probability.

Subjects were instructed to fixate centrally throughout the task and to respond as fast as possible to two targets, the letters “L” and “T”, with left and right button-presses respectively. The experimenter reminded subjects of instructions periodically. Eye movements of three of six subjects were tracked using the EyeLink 1000 (SR Research, Ottawa, Ontario, Canada) in order to verify central gaze fixation in a previous study⁹³. Eye tracking for all subjects was not possible due to interference caused by bandages covering regions surrounding the eyes. The experimenter watched subjects and noted trials with excess movement or breaks in visual fixation, so they could be removed from analyses. Additional recordings taken prior to the task, at the start of each recording session, served as a baseline period. We focused our study on how PAC during the cue period relates to spatially and behaviorally defined sites. We define the cue period as the period between the onset of the cue and the onset of the target. Sites were functionally classified by neural activity during the target period, which is defined as the first 400 ms after the target appears.

In the target period we classified cortical sites as “spatial” or “behavioral” based on local neural activity that discriminated spatial task variables or behavioral responses. Spatial sites had high-gamma power that discriminated target location (i.e. contralateral vs ipsilateral to recording sites) or target validity (i.e. valid vs invalid). We classified behavioral sites based on significant Spearman correlation between high-gamma power in the target period and RT. We removed cortical sites with both spatial and behavioral classifications from further analysis due to their limited number.

Digital Signal Processing

We performed all digital signal analysis with custom scripts in MATLAB (The MathWorks Inc, Natick, MA). A custom graphical user interface was developed to visually inspect temporal and spectral properties of every channel. Channels with abnormal amplitude (e.g. $> \pm 1000$ mV) or power spectra (e.g. harmonic noise) were flagged. Time periods containing transient artifacts across groups of channels were flagged. All flagged channels and time periods were not used in further analysis. We performed spectral decomposition using Morlet wavelet convolution and estimated phase and amplitude envelopes from the resulting complex signals. All signals were then filtered and down-sampled to 300 Hz. All wavelet-derived properties (i.e. phase, amplitude and power) are generated from the whole signal, before trials are extracted, to avoid edge effects.

Two sets of wavelet libraries were used for phase amplitude coupling. We created these libraries to satisfy mathematical constraints on phase-amplitude coupling measurements. Specifically, the bandwidth of the frequency-for-amplitude (Fa) must be twice the frequency-for-phase (Fp) of interest⁷⁴. The two wavelet libraries were constructed as follows.

Frequency for amplitude wavelets: We used the full width at half-maximum (FWHM) of the Morlet wavelet as a lower bound estimate for bandwidth. We designed Fa wavelets to have a FWHM of 20 Hz and used 21 wavelets with center frequencies ranging from 20 Hz to 150 Hz in 5Hz increments.

Frequency for phase wavelets: We designed narrow-band Fp wavelets for phase specificity. Higher frequency resolution was employed for phase signals to distinguish between delta, theta and alpha rhythms. We used 20 Fp wavelets ranging from 1 Hz to 20 Hz with 1Hz spacing and FWHM of 0.8 Hz.

Quantifying Phase-Amplitude Coupling with the Modulation Index

We measured PAC using the modulation index (MI) ⁷³, which quantifies the magnitude of coupling. MI also provides a common measurement to compare different forms of PAC (e.g. unimodal vs bimodal) across different frequencies. MI was calculated as the Kullback-Leibler divergence between the uniform distribution (i.e. pure entropy) and the observed probability density $P(j)$, which describes the normalized mean amplitude at a given binned phase (see $P(j)$ below). Pairwise calculation of MIs for two sequences of frequencies produces a comodulogram. MI is calculated as follows:

$$MI = \frac{D_{KL}(P, Q)}{\log(N)}$$

$$D_{KL}(P, Q) = \sum_{j=1}^N P(j) \log\left(\frac{P(j)}{Q(j)}\right)$$

Where D_{KL} is the Kullback-Leibler divergence, P is the observed phase-amplitude probability density function, Q is the uniform distribution and N is the number of phase bins. P follows the equation:

$$P(j) = \frac{\langle A_{f_A} \rangle_{\phi_{f_P}}(j)}{\sum_{k=1}^N \langle A_{f_A} \rangle_{\phi_{f_P}}(k)}$$

where $\langle A_{f_A} \rangle_{\phi_{f_P}}(j)$ is the mean f_A amplitude signal at phase bin j of the phase signal ϕ_{f_P} . We divided phase into 18 bins of 20-degree intervals. For a review of PAC methods refer to ⁷³.

To identify PAC frequency pairs of interest, we sorted trials by RT and divided them into quartiles (**Figure 2 e-g**). We used signals from the fastest and slowest quartiles to generate $P(j)$ distributions of normalized amplitude per binned phase, from which we calculated the MI. We

verified the precision of our methods with simulations using methods defined in the appendix of Tort et al.⁷³(**Supplemental Figure 1 b-f**). Specific parameters or MATLAB scripts used for simulations are available upon request.

Statistical Analysis

Band Limited Power and PAC Time Series Comparisons: Statistical inference testing of band-limited power and PAC time series followed methods described by Maris and Oostenveld⁹². Cluster candidates were generated using t-statistics to test the null hypothesis that there was no difference between categories at each sample. If a sample t-statistic exceeded an alpha level of 5% then the null hypothesis was rejected for the sample and it was considered a cluster candidate. Temporally adjacent cluster candidates were grouped into a single cluster and their t-statistics were summed to produce a clustering statistic. The clustering statistic of the observed data was tested against a permutation distribution. To produce the permutation distribution, trial labels (e.g. valid vs invalid) are shuffled and randomly reassigned 10,000 times. For each shuffle, cluster candidates and clustering statistics were generated as described above. The maximum clustering statistic from each shuffle was used to create the permutation distribution. We calculated p-values for observed clusters using the formula $p = (r+1)/(n+1)$, where r is the number of shuffled clustering statistics greater than the observed clustering statistic and n is the total number of shuffled sets used⁹⁵. We corrected for multiple comparisons across cortical sites with the False Discovery Rate (FDR) correction method.

Phase-Amplitude Coupling Comparison: We adapted a two-dimensional non-parametric permutation test to make cluster-based statistical inferences on comodulograms based on the difference between fast and slow trials. First, we generated 1,500 shuffled distributions for each

cortical site by randomly reassigning RTs to trials, sorting, dividing into quartiles, and calculating the absolute difference in comodulograms for fast and slow trial quartiles as follows:

$$d_{fAfP} = |MI_{fAfP}^{fast} - MI_{fAfP}^{slow}|$$

We use the pooled variance in each frequency pair in the distribution of $d_{fAfP}^{shuffled}$ to determine the cutoff threshold specific to each frequency pair. Adjacent supra-threshold frequency-pairs were grouped together in clusters and t-statistics were summed. We tested the null hypothesis that the shuffled data was no different from the observed data using a two-dimensional cluster based permutation test where diagonals were not considered neighbors ⁹².

PAC time series were calculated using MI calculations in a 500 ms sliding window with 50 ms increments. While this window only includes half a 1 Hz cycle, we empirically confirmed that the large amount of data (>250 seconds) used in these analyses ensured that all phases of the 1 Hz cycle were represented in the MI calculation. Differences between PAC time-series for spatial and behavioral site categories were calculated with the one-dimensional cluster-based permutation test described above.

Inter-trial coherence and preferred phase statistics: Inter-trial coherence is the magnitude of the mean phase across trials. It reflects the phase consistency across trials for every time point and frequency. Preferred phases were calculated as the maximum phase-bin in the phase-amplitude probability density plot (see $P(j)$ above). Preferred phases were calculated separately for each cortical site. The non-uniformity of preferred phases was determined with the Rayleigh test and the equivalence of the circular means for spatial and behavioral sites was calculated with the Kuiper test ⁹⁶.

Sharp Waveform Detection

To detect the presence of sharp waves we employed methods from QRS detection in ECG analyses^{97,98}. We used the first differential of the ECoG Signal to identify periods of rapid change. The Hilbert transformation is then applied as an envelope function where peaks corresponded to locations of candidate sharp wave. We set a threshold of one standard deviation from the mean. Any candidate sharp wave that did not surpass this threshold for more than 16 ms was rejected. Finally, we calculated the amplitude change, or height, of each candidate sharp wave and rejected the bottom 80% to ensure that the most prominent waveforms were being isolated.

4.4 Results

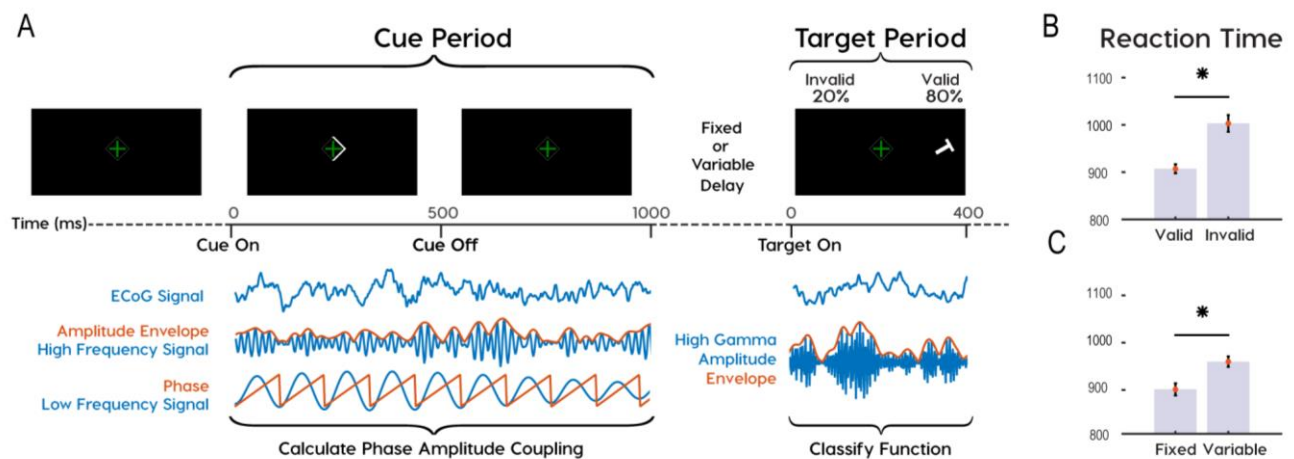


Figure 4.4.1 Task design showing analysis periods and behavioral results: (A) Subjects participated in a Posner cued attention task, with the most probably target location indicated by an arrowhead presented at fixation prior to target onset. Phase-amplitude coupling (PAC) was analyzed during the cue period. Functional classifications were based on high-gamma amplitude during the target period. Subjects responded to “L” and “T” targets with left and right mouse clicks. (B) Reaction times were greater for invalid than valid trials and (C) for variable than fixed delays.

Reaction time reflects task performance

We employed a spatial cueing task to induce and measure covert shifts in spatial attention. Participants fixated on a central crosshair throughout the task. At the beginning of each trial a central arrow, or “cue”, pointed towards the most likely location of a subsequent “target”. Two target locations were possible. We defined these locations as “contralateral” or “ipsilateral”, depending on whether it was on the same side, or the opposite side, of the recording sites. The cue predicted the location of the target in 80% of trials (i.e. valid trials). In 20% of trials the target appeared opposite to where the cue pointed (i.e. invalid trials). In our version of the task participants were required to discriminate “L” from “T” targets that were randomly rotated (**Figure 4.4.1, top**). Across six participants, 5100 of 5344 trials were completed with 92.2% correct responses. We used trials with correct responses for further analyses. Invalid trials incurred higher RTs than valid trials (medians (ms): valid = 831, invalid = 927, rank sum: 7.2e6, approximate z-value: -9.8, $p = 1.6e-22$). No differences in reaction time occurred between trials with contralateral and ipsilateral cues (medians (ms): contralateral=860, ipsilateral=863, rank sum: 5.1e6, approximate z-value: -1.7, $p=0.08$). Higher RTs on invalid trials confirmed that the task induced lateralized shifts in visual attention.

We focused our physiological analyses on two periods in the task, the “cue period” and the “target period”. We define the “cue period” as the time between cue onset and target onset and the “target period” as the first 400ms after target onset. Cued shifts in attention occurred during the cue period. The target location and identity is revealed during the target period, allowing subjects to locate the target, identify it, and subsequently respond appropriately.

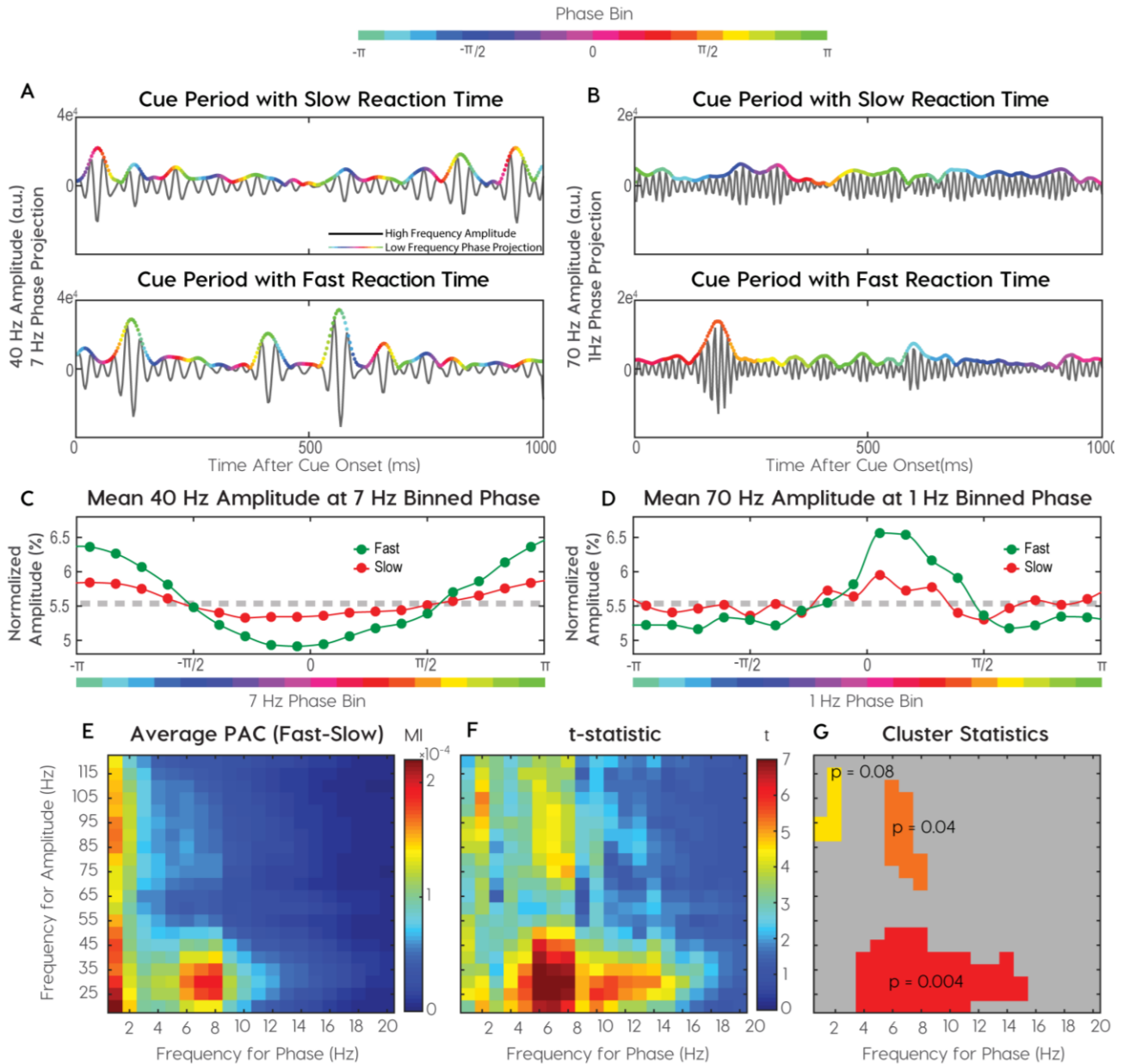


Figure 4.4.2 Identification of distinct phase-amplitude coupling clusters: (A-B) Single trial

examples of more coupling on fast trials than slow trials (A) Beta (25 Hz) amplitude

couples to alpha phase (8 Hz) and (B) gamma (70 Hz) amplitude couples to delta (1 Hz) phase. High-frequency amplitude envelopes are colored with low-frequency phase. (C) Mean beta amplitude at binned alpha phase and (D) mean gamma amplitude at binned delta phase. € Mean modulation indices calculated for pairwise frequencies. (F) T-statistics from permutation distributions were used to determine significance. (G) Delta-high gamma (DH, yellow), theta/alpha-beta/low gamma (TABL, red) and theta/alpha-high gamma (orange) clusters, calculated during the cue period predicted reaction time.

Distinct clusters of PAC predict RT:

We identified behaviorally relevant PAC frequencies by comparing the cue periods preceding fast and slow RTs. After sorting trials by RT, we binned them into quartiles and analyzed differences in cue period PAC between the fastest and slowest quartiles. We found PAC differences in multiple frequency pairs (**Figure 4.4.2a, b**). For visualization purposes, color-coded low-frequency phases are projected on the high-frequency amplitude envelope (for reference see **Supplemental Figure 1 b, d**). The modulation index (MI) quantified PAC intensity, or the non-uniformity of the phase-amplitude probability density plot (**Figure 4.4.2c, d**). We calculate the MI for each frequency pair to generate a comodulogram. The absolute difference between slow and fast comodulograms averaged over all cortical sites and subjects, revealed coupling between delta-phase and theta-alpha-phase to higher frequency (>20Hz) amplitude envelopes (**Figure 4.4.2e**).

Cluster-based permutation testing determined the significance of PAC while controlling for false positives. Permutation distributions were generated by shuffling reaction times between trials and calculating absolute differences in comodulograms between the slowest and fastest quartiles in shuffled datasets. Significant phase-amplitude frequency clusters outlined adjacent

frequency pairs with significant t-statistics (**Figure 4.4.2f**). Three clusters of frequency pairings explained RT variance: 1) theta-alpha-phase/beta-low gamma-amplitude (TABL) (**Figure 4.4.2g, red**), 2) delta-phase/high-gamma-amplitude (DH) (**Figure 4.4.2g, yellow**), and 3) alpha-phase/high-gamma-amplitude (**Figure 4.4.2g, orange**). We focused on comparing DH with TABL PAC for three reasons. First, they have unique phase and amplitude frequencies, unlike alpha-high gamma PAC. Second, the magnitudes of DH and TABL PAC were the least correlated across all electrodes. Third, TABL and DH PAC have been shown to underlie distinct mechanisms in human memory⁶⁹. Therefore, we hypothesized that DH and TABL were most likely to show different functional characteristics during the spatial attention task.

To verify precision in measuring frequency-specific PAC we modeled PAC with simulated signals using procedures from Tort and colleagues⁷³ (**Supplemental Figure 1 a-e**). We inspected subject-specific differences in PAC by inspecting averaged comodulograms alongside electrode locations on an averaged brain plots. Five of six subjects displayed robust TABL PAC. Most subjects had less DH PAC magnitude compared to TABL PAC (**Supplemental Figure 3**). Interestingly, TABL PAC did not appear on single trials and only emerged when we calculated phase-amplitude distributions on multiple (>15) trials (**Supplemental Figure 2**). To explore differences in TABL and DH PAC we returned to disparities in the motivating literature. While both Esghaei et al. and Szczepanski et al. found that PAC depended on spatial task variables, only the latter found that PAC correlated with RT^{48,75}. Therefore, we explored whether DH and TABL PAC uniquely depended on spatial variables or task performance (i.e. RT) by classifying cortical sites by function and comparing PAC across functional classes.

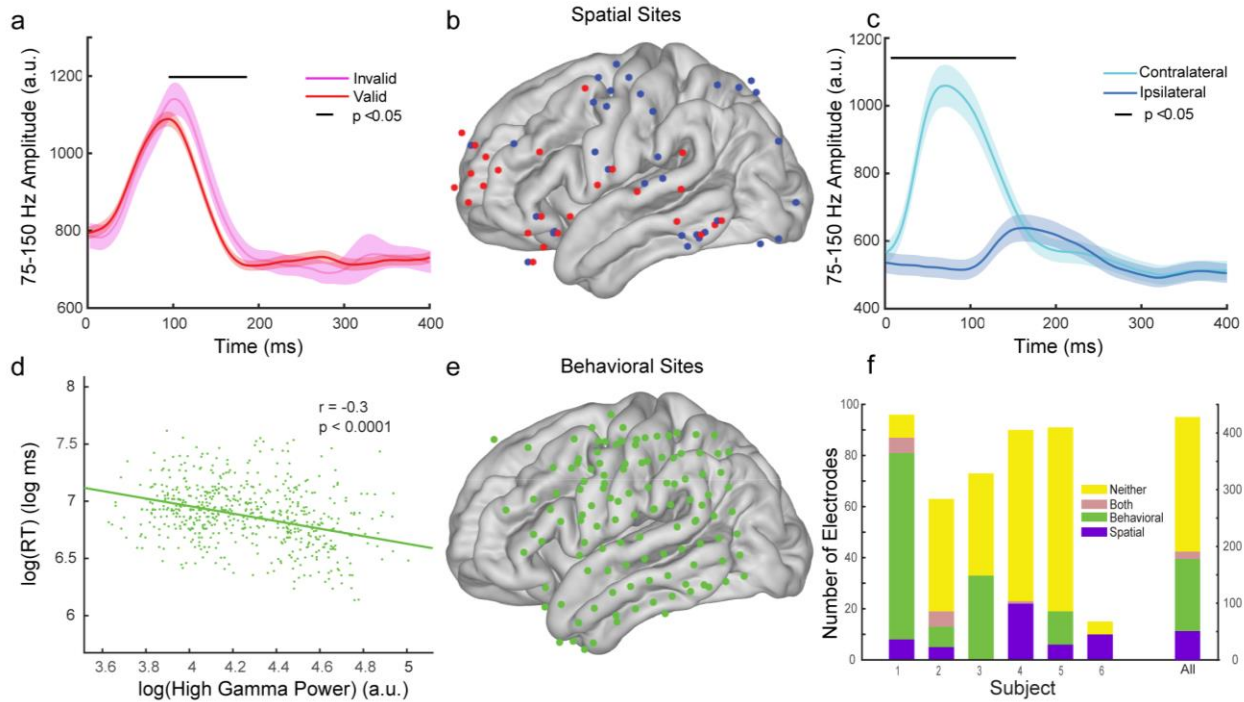


Figure 4.4.3: Functional classification of “spatial and “behavioral” cortical sites. Spatial

cortical sites have high gamma (75-150 Hz) activity that discriminates spatial parameters of the task. Exemplary mean high gamma power, with standard error bars, from two spatial cortical sites discriminating (A) valide from invalid targets and (C) targets that appear contralateral from targets that appear ipsilateral to recording sites. (B) Distribution of spatial cortical sites where red dots discriminate validity and blue dots discriminate location. Behavioral cortical sites had high gamma activity that correlated with reaction times (RTs). (D) exemplary correlation from one behavioral cortical site. (E) The cortical surface distribution of behavioral sites. (F) Breakdown of electrode classifications for each participant (left vertical axis) and for all participants (right vertical axis)

Functional classification of spatial and behavioral cortical sites:

We classified cortical sites based on high-gamma (75-150 Hz) power as it has been shown to correlate with local neuronal activity⁹⁹. We classified neural activity in the target period for two reasons. First, this period had the most robust discriminatory neural responses. Second, all the information required for a correct response was made available in the target period. Therefore, all classifications relied on high-gamma power in the first 400 milliseconds after target onset.

We defined "spatial" cortical sites based on neural activity discriminating target location or validity. Due to the contralateral organization of visual receptive fields, contralateral targets typically produced larger neural responses than ipsilateral targets. Invalid targets, occurring on 20% of trials, typically produced larger changes in high-gamma power than valid targets. We justified grouping validity sensitive sites with location sensitive sites because calculating a target's validity requires the target's location as an operand. Thus, we classified a cortical site as "spatial" when its high-gamma power discriminated either contralateral from ipsilateral targets (**Figure 4.4.3a**) or valid from invalid targets (**Figure 4.4.3c**, cluster permutation test threshold $p < 0.05$, FDR corrected). In contrast, we defined "behavioral" cortical sites based on discriminatory neural activity pertaining to RT. Behavioral cortical sites had significant Spearman correlations between mean post-target high-gamma power and RT (Spearman correlation threshold $p < 0.05$, FDR corrected). Mean high-gamma and RT had log-normal distributions so we represent this as a correlation of the log of both values in **Figure 4.4.3d**. Spurious correlations were ruled out with a permutation test using 5000 identical correlations with shuffled RTs.

We found 51 spatial and 127 behavioral sites. We removed the 13 sites classified to both groups and 237 classified to neither group, from further analyses (**Figure 4.4.3f**). Spatial cortical sites were found primarily over visual cortex, parietal cortex, motor cortex and frontal cortex, with the distribution of dorsal parietal and posterior frontal sites roughly matching the dorsal attention network¹¹ (**Figure 4.4.3b**). Behavioral cortical sites were found over motor cortex, parietal cortex, and temporal cortex (**Figure 4.4.3e**). We compared these two functional classes to investigate functional properties of TABL and DH PAC.

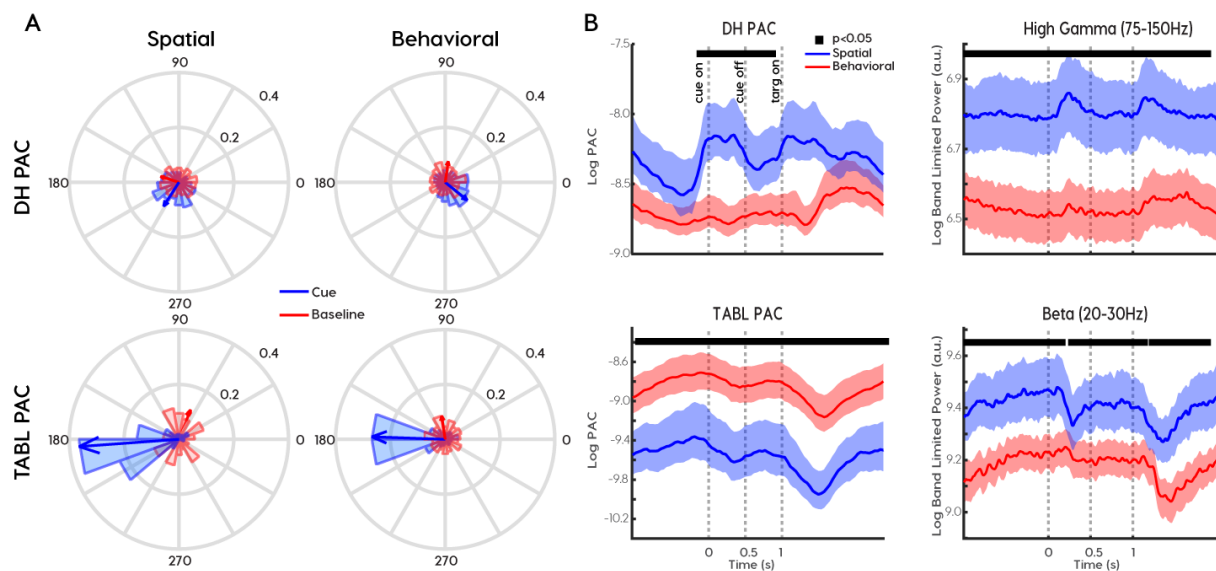


Figure 4.4.4 Phase preferences and relative PAC magnitude distinguish coupling frequencies and functional classes. (A) Preferred phases differed for delta to high-gamma (DH) PAC (top row) and theta-alpha to beta-low-gamma TABL PAC (bottom row) across spatial (left column) and behavioral (right column) cortical sites. Coupling emerged in the cue period (blue) relative to baseline pseudo-trials (red). Arrows indicate the circular mean of preferred phases across all subjects and cortical sites. (B) The relationship between PAC magnitude (left column) and the corresponding frequency-for-amplitude power (right column) reverses across behavioral (red) and spatial (blue) cortical sites. Solid black lines

at top of graph indicates clusters of samples where spatial and behavioral signals were significantly different at $p < 0.05$.

Phase and magnitude differentiate PAC clusters across functional classes:

We first characterized differences in the phase preferences and coupling magnitudes of DH and TABL PAC. We defined preferred phases as the low-frequency phase bin with the greatest high-frequency amplitude. Preferred phases are independent of coupling magnitude, and if no coupling exists then preferred phases will be uniformly distributed on average. We calculated the preferred phase using all cue periods, then calculated the circular mean across all cortical sites within a functional class. We repeated this analysis for pseudo-trials generated from the rest period recorded prior to the task to ensure that coupling did not exist during rest. Pseudo-trials had identical length to the cue period, but were sampled randomly from the rest period.

During rest, preferred phases were uniformly distributed in both functional classes and frequency pairs (Rayleigh test, TABL PAC: spatial $z = 1.5$, $p = 0.86$; behavioral $z = 0.97$, $p = 0.38$; DH PAC: spatial $z = 0.03$, $p = 0.97$; behavioral $z = 2.86$, $p = 0.06$; **Figure 4.4.4A, red**). In contrast, all sites had preferred phases during the cue period. (Rayleigh test, TABL PAC: spatial $z = 28$, $p = 1e-12$; behavioral $z = 41$, $p = 6e-20$; DH PAC: spatial $z = 37$, $p = 3e-9$; behavioral $z = 71$, $p = 4e-32$, **Figure 4.4.4A, blue**). Furthermore, the mean DH PAC phase preferences differed across spatial and behavioral sites (spatial mean=237, std=67, behavioral mean=321, std=69, Kuiper test, $k = 1.1e5$, $p = 0.001$, **Figure 4.4.4A top**). However, TABL PAC consistently showed a 180-degree phase preference in all sites (spatial mean=184, std=42, behavioral

mean=178, std=53; Kuiper test, $k = 1.7e4$, $p = 0.10$, **Figure 4.4.4A bottom**). Phase preferences were not affected by cue direction or reaction time. Therefore, the preferred phases of DH and TABL PAC during the cue period showed consistent differences.

We hypothesized that spatial sites were more active than behavioral sites during the cue period. To verify this, we compared high-gamma variance and inter-trial coherence (ITC) across functional classes. High-gamma power and variance were greater in spatial sites than in behavioral sites (power: cluster permutation test $p < 0.001$ **Figure 4.4.4B, top right**, variance: rank sum, $p < 0.001$ **Supplementary Figure 4A**). ITC measures the coherence over trials in both phase and amplitude. Again, spatial sites had greater ITC in the 1-10 Hz frequency range during the cue period (**Supplemental Figure 4B, C**). Taken together, neural activity in spatial sites were more modulated by spatial cueing than behavioral sites were in both low and high frequencies. DH PAC paralleled these indexes of neural modulation and was higher in spatial sites than in behavioral sites (cluster permutation test, $p < 0.001$, **Figure 4.4.4B top left**). Furthermore, DH PAC peaked after cue onset and target onset, much like high gamma power. In contrast, TABL PAC was greater in behavioral sites than in spatial sites (cluster permutation test, $p < 0.001$, **Figure 4.4.4B bottom left**). This was not due to high beta or low-gamma power, both of which were greater in spatial sites (cluster permutation test, $p < 0.001$, **Figure 4.4.4B bottom right**). It was also not due to greater alpha activity in behavioral sites (**Supplemental Figure 4E**). Finally, this was not due to baseline differences as there were no significant differences in beta, low-gamma or alpha power at rest across the functional classes. Therefore, high TABL PAC corresponded to reduced neural modulation across the functional categories.

To summarize, TABL PAC and DH PAC showed a reversal in their relationship to power. TABL PAC was greater in behavioral sites, despite reduced power, variance and ITC. In

contrast, DH PAC positively co-varied with every measure of neural modulation we measured. Since there were no observed differences in PAC or power at baseline, this reversal was best explained by the task. Along with differences in phase preference, magnitude differences provided additional evidence that DH and TABL PAC dynamics depended on functional classifications.

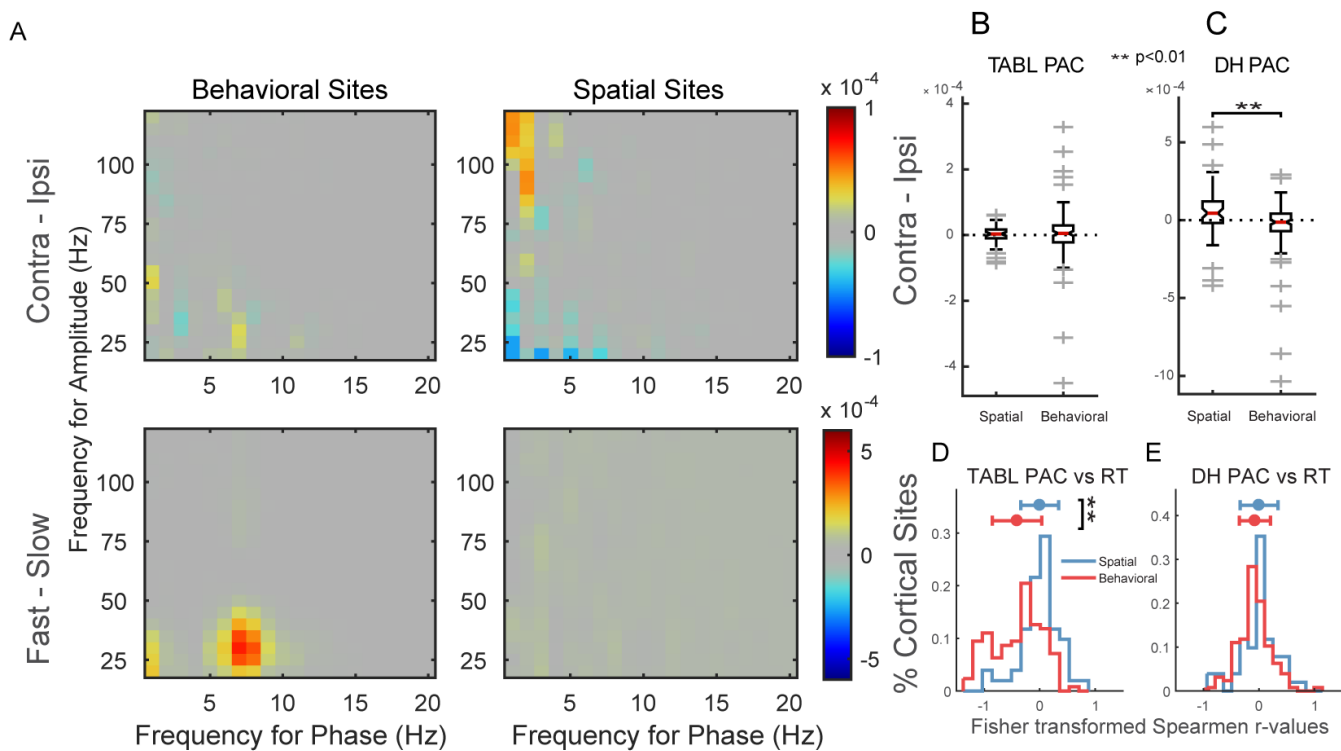


Figure 4.4.5 PAC magnitude differences across task conditions distinguish between phase frequencies and functional classes. (A) Shows positive comodulogram differences for behavioral (left) and spatial (Right) sites when subtracting contralateral from ipsilateral trials (top) and fast from slow trials (bottom). These changes were quantified using previously identified PAC clusters. (B) No ipsi-contra differences were observed as a product of lateralized cued attention in theta/alpha-beta/low gamma (TABL) PAC at either

spatial or behavioral sites. (C) Delta-high gamma (DH) PAC increases on contralateral vs ipsilateral trials in spatial sites relative to behavioral sites. (D) Table PAC was more negatively correlated with RT in behavioral than spatial sites. DH PAC showed no differences between behavioral and spatial sites (E).

Distinct PAC frequency pairs correspond to cortical function

We investigated PAC differences across spatial conditions (i.e. contralateral vs ipsilateral cues) and behavioral responses (i.e. RT) (**Figure 4.4.5A**). We quantified differences using previously defined clusters (**Figure 4.4.2G**). On contralateral trials, we found increased DH PAC in spatial cortical sites and no change in behavioral sites (sign test, spatial: z-value 2.6, $p = 0.009$, behavioral: z-value = -1.8, $p = 0.07$; **Figure 4.4.5C**). The difference between classes was significant (rank sum, z-value = -3.4, $p = 7e-4$, corrected). TABL PAC showed no changes across ipsilateral and contralateral cueing conditions and there were no differences between classes (sign test, spatial: z-value 1.05, $p = 0.29$, behavioral: z-value = 0.94, $p = 0.34$, rank sum, z-value = 0.15, $p = 0.9$, **Figure 4.4.5B**). This suggested DH PAC was specific to spatial variables in spatial sites.

To understand which functional class drove differences between fast and slow trials we correlated DH and TABLPAC with RT for all sites in each functional class. Spatial sites showed no significant trend in correlations (t-test, DH vs RT $\neq 0$: $p = 0.80$, TABL vs RT $\neq 0$: $p = 0.97$) and there were no differences between Fisher transformed r-value distributions (t-test, DH vs RT \neq TABL vs RT: $p=0.90$, **Figure 4.4.5E**). In contrast, TABL PAC was more negatively correlated with RT in behavioral sites (t-test, DH vs RT \neq TABL vs RT: $p<1e-11$, **Figure 4.4.5D**). To

summarize, during the cue period DH PAC in spatial sites varied with cue direction and TABL PAC in behavioral sites correlated with RT.

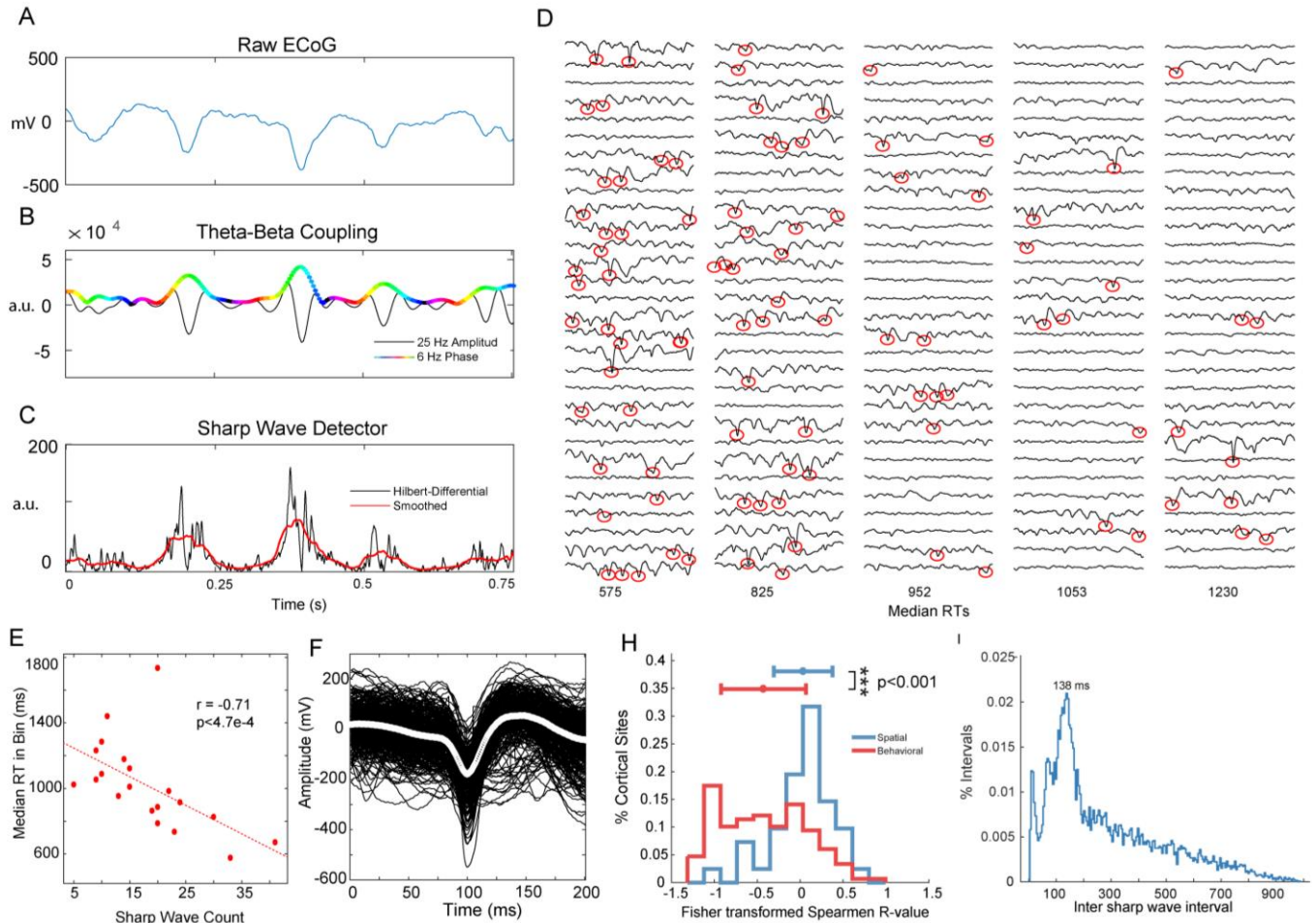


Figure 4.4.6: Transient waves are behaviorally relevant and coupled to 7.2 Hz phase. (A) Raw ECoG signal shows three sharp waves that correspond to (B) theta (6 Hz)-phase, beta (25 Hz)-amplitude coupling with a 180-degree phase preference. (C) Hilbert transform of the differential of the raw ECoG signal amplifies rapid (i.e. sharp) changes in the ECoG signal. A threshold on the Hilbert-Differential identified the transient waves. (D) Single trials sorted by RT and grouped into 30-trial bins with median RTs below each column. Red circles highlight identified sharp waves that were more likely to occur on trials with

fast reaction times. € Correlation between the number of sharp waves in an RT-bin and RT. (F) the averaged waveform for the cortical site in (D) and (E). (H) the distribution of correlation coefficients for spatial and behavioral sites. (I) distribution of inter-sharp wave intervals peak at 138 ms, corresponding to 7.2 Hz.

Coupled sharp waves correspond to TABL PAC:

DH PAC has been described in prior attention literature^{45,48,75}, but TABL PAC has not been previously shown in attention tasks. It has, however, been shown to correlate with encoding and retrieval in memory tasks⁶⁹. Additionally, non-sinusoidal or “sharp” waveforms have been shown to produce coupling in the TABL PAC frequency range^{69,86}. We hypothesized that if sharp waveforms resulted in TABL PAC, then they too will correlate with RT. Furthermore, we hypothesized that, compared to computationally expensive PAC analyses, more efficient methods could be used to detect and characterize sharp waveforms.

On visual inspection of the raw ECoG we found that transient waveforms temporally co-occurred with TABL PAC (**Figure 4.4.6A, B**). We reasoned that the “sharpness” of the wave could be used to detect these events and employed techniques previously used for QRS wave detection in electrocardiogram analyses^{97,98}. This method uses the Hilbert transform of the differential to identify rapid (i.e. sharp) changes in the raw signal (**Figure 4.4.6C**). We applied this sharp wave detection to individual trials and displayed the results in 30-trial RT bins (**Figure 4.4.6D**). We found that trials with fast responses (i.e. low RTs) were more likely to have sharp waveforms than slower trials (**Figure 4.4.6E**). In aggregate, sharp waveform correlations with RT were significantly more negative in behavioral sites than in spatial sites (rank sum $p < 0.001$,

Figure 4.4.6H), similarly to TABL PAC (**Figure 4.4.5D**). Alongside the correlations for specific channels we used the minimum of the detected sharp wave to time-lock waveforms and observe their shapes. We observed that detected sharp waves consisted of a roughly 50 ms transient wave (**Figure 4.4.6F**).

Finally, it remained unclear whether these sharp waves were themselves coupled to a low frequency rhythm, so we analyzed the intervals between sharp waves. For every trial with more than one sharp wave we recorded the interval between adjacent sharp waves. The distribution of inter-sharp-wave intervals was not log-normal or Poisson (chi-square goodness-of-fit both $p < 0.0001$), which would be expected if intervals were randomly spaced. Instead, a peak of 138 ms suggested that sharp waves coupled to a 7.2 Hz oscillation (**Figure 4.4.6I**).

In summary, TABL PAC associated with sharp waveforms that could be detected with time-domain methods. However, the distribution of sharp wave intervals suggested that they occur regularly within a theta-alpha range oscillation. Therefore, sharp wave morphometry and the frequency of sharp waveform events contributes to TABL PAC. While characterization of these transient waveforms may require nuance, their correlations with RT across functional classes matched that of TABL PAC, suggesting a common origin.

4.5 Discussion

This study expands our understanding of phase amplitude coupling (PAC) in several ways. First, we identified an interaction between theta-alpha-phase and beta-low-gamma amplitude (TABL PAC) during spatial cueing that predicted future reaction times (RTs) and is novel to the attention literature. Second, we characterized differences between TABL PAC and delta-phase, high-gamma-amplitude coupling (DH) PAC. Specifically, we found DH PAC

positively covaried with contralaterally cued attention in spatial sites while TABL PAC correlated with RT in behavioral sites. Unlike DH PAC, TABL PAC negatively covaried with multiple indices of neural modulation and maintained a consistent 180-degree preferred phase during the task. Third, we identified sharp wave correlates of TABL PAC using novel and efficient methods, which showed that sharp waveforms predict RT. Finally, we showed that behaviorally relevant sharp waveforms are also coupled with low frequency oscillations. Taken together, this study highlights the unique contributions of distinct PAC frequency clusters in a spatial attention task.

Methodological considerations for frequency-specificity in PAC analysis:

The complexity of PAC analyses⁷⁴, evidence for spurious PAC^{87,100}, and the novelty of TABL PAC in spatial attention tasks warrants a discussion of methods. We employed the modulation index (MI) to measure the intensity of PAC due to its advantages over other methods (for a review, see Tort et al., 2010). Sufficient data length was critical to resolving TABL PAC. This was due to the relative sparsity of transient waveforms leading to TABL PAC on single trials. Additionally, coupling between two frequencies (e.g. Fp and Fa) can only occur in the presence of two additional frequencies: the sum and the difference of the original two frequencies (i.e. Fa+Fp and Fa-Fp, see **Supplemental Figure 1A arrows**). To account for this, we developed wavelet libraries independently for phase and amplitude signals. Wide band-width amplitude filters were required to measure PAC in higher phases^{74,101}. Conversely, narrow band-widths allowed phase-specificity when filtering phase signals, since broadband phase estimates may include separable PAC components that cancel each other out.

In addition to signal processing and measurement considerations, we used reaction time to identify behaviorally relevant PAC frequency clusters. Furthermore, instead of shuffling

phase and amplitude signals relative to each other, we shuffled trials between behavioral categories (e.g. fast and slow RT quartiles) to generate chance distributions. This guaranteed that our statistical inferences were made from distributions with physiologically observed phase-amplitude relationships. We then employed a non-parametric cluster-based permutation test for statistical inferences. These methods resolved behaviorally relevant PAC frequency clusters spanning frequency bands previously associated with attentional processes ^{42,44,46,67}

Sharp Waveforms are Dynamically Coupled

We propose that sharp waveforms are a behaviorally relevant, special case of PAC. This is an alternative to suggestions that non-sinusoidal sharp waveforms are spurious ^{86,102}. We demonstrated transient beta (50 ms) waves that occurred at 138 ms intervals, suggesting a coupled dynamical process. Similar waveforms have been shown across species ¹⁰³ and have relevance to human memory tasks ⁶⁹. Sharp waveforms have high frequency components coupled to the phase of low frequency rhythms, which fits the definition of PAC. In the context of dynamic brain activity, persistent nesting of sinusoidal oscillations is another special case of PAC, not the only possibility. PAC approaches identify frequency interactions between high and low frequencies that can lead to a better understanding of underlying dynamics. Because PAC analyses are computationally expensive and sensitive to noise, we used our PAC approach to inform a computationally inexpensive approach, which better described individual trials.

A theoretical framework and implications for PAC in attention

Two hypotheses are relevant to PAC and cued spatial attention (for an extensive review see ⁸³. The communication through coherence (CTC) hypothesis proposes that excitability

windows of two communicating brain regions are temporally aligned by low frequency oscillations to promote information transfer (e.g. spikes) across regions^{104,105}. In contrast, the gating by inhibition (GBI) hypothesis proposes that low frequencies periodically suppress information in a neuronal population¹⁰⁶.

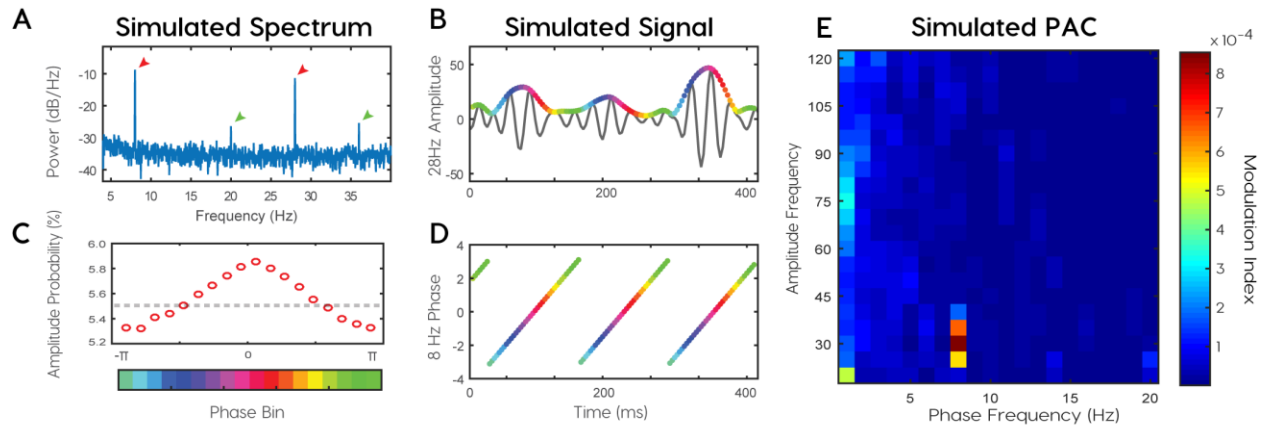
Although we do not have evidence to assess the CTC hypothesis, we found evidence for a suppressive role for TABL PAC, which fits with the GBI hypothesis. First, multiple indices of neural modulation suggested spatial sites were more active than behavioral sites during spatial cueing, yet TABL PAC was greater in behavioral than spatial sites. Furthermore, TABL PAC consistently demonstrated a 180-degree phase preference, which meant that high frequency activity preferred the trough of the low frequency oscillation. This pattern of coupling has been shown in “pulsed-inhibition” or “gating by inhibition” models of alpha oscillatory activity^{47,106}, where the peak of alpha oscillatory amplitude periodically suppresses higher frequency activity. Finally, the transient waves that cause TABL PAC have been shown in multiple species and studies to relate to decreased information relay^{103,107}. Taken together these results support a suppressive function of TABL PAC during our spatial attention task.

Conclusions

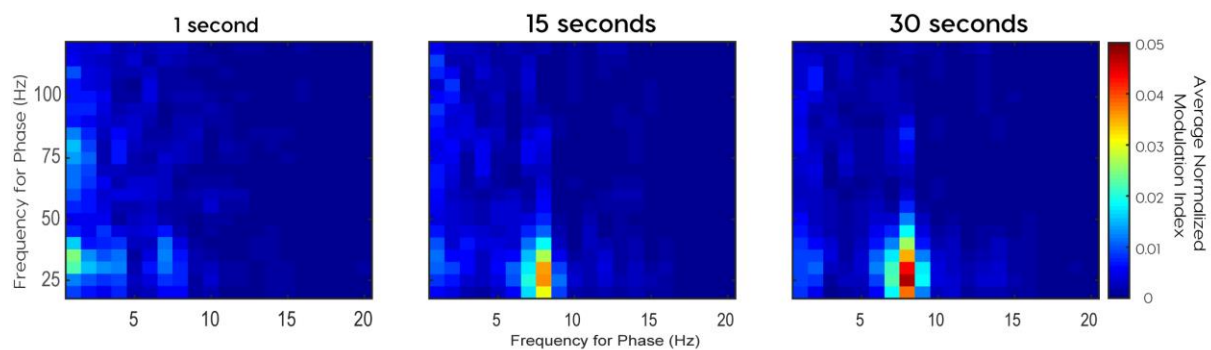
Spatial attention enhances neural responses to attended stimuli and by filtering out unwanted information¹⁰⁸. Prior studies differed on whether PAC contributes to selection⁴⁸ or filtering⁷⁵. Here we show evidence that the functional properties of PAC depend critically on phase frequency. DH PAC facilitates lateralized spatial attention while TABL PAC contributes to behavioral performance, potentially through suppressive gating of activity. Furthermore, we demonstrate that sharp waves producing TABL PAC are themselves coupled to low frequency

oscillations. While the function of these sharp waves remains unclear, PAC remains a useful tool in understanding the neural correlates of human attention.

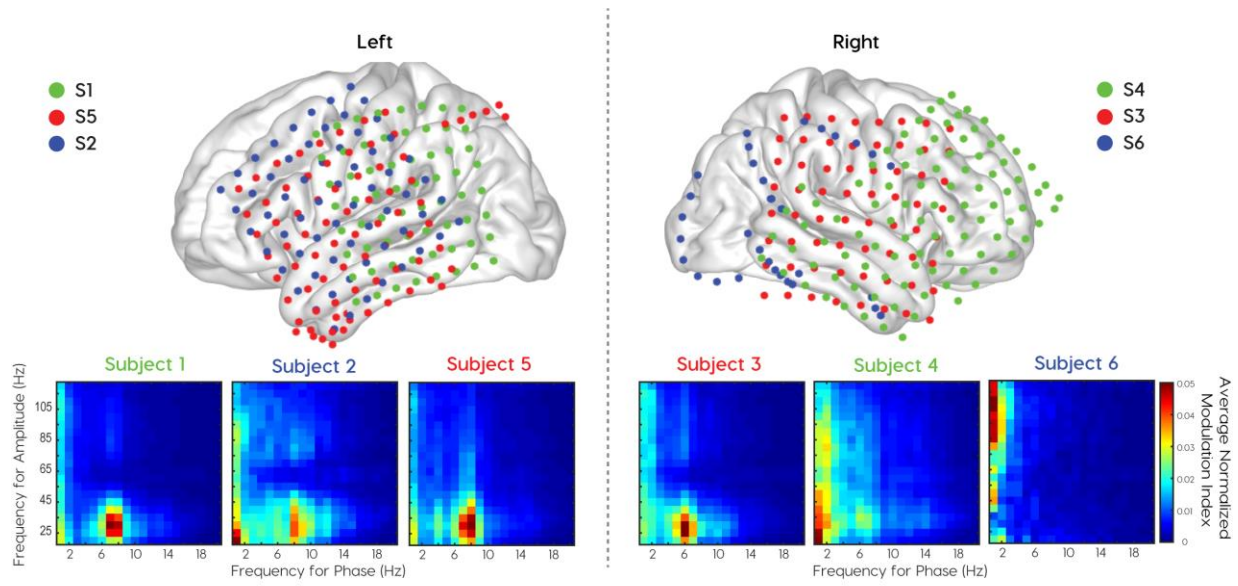
4.6 Supplemental Figures:



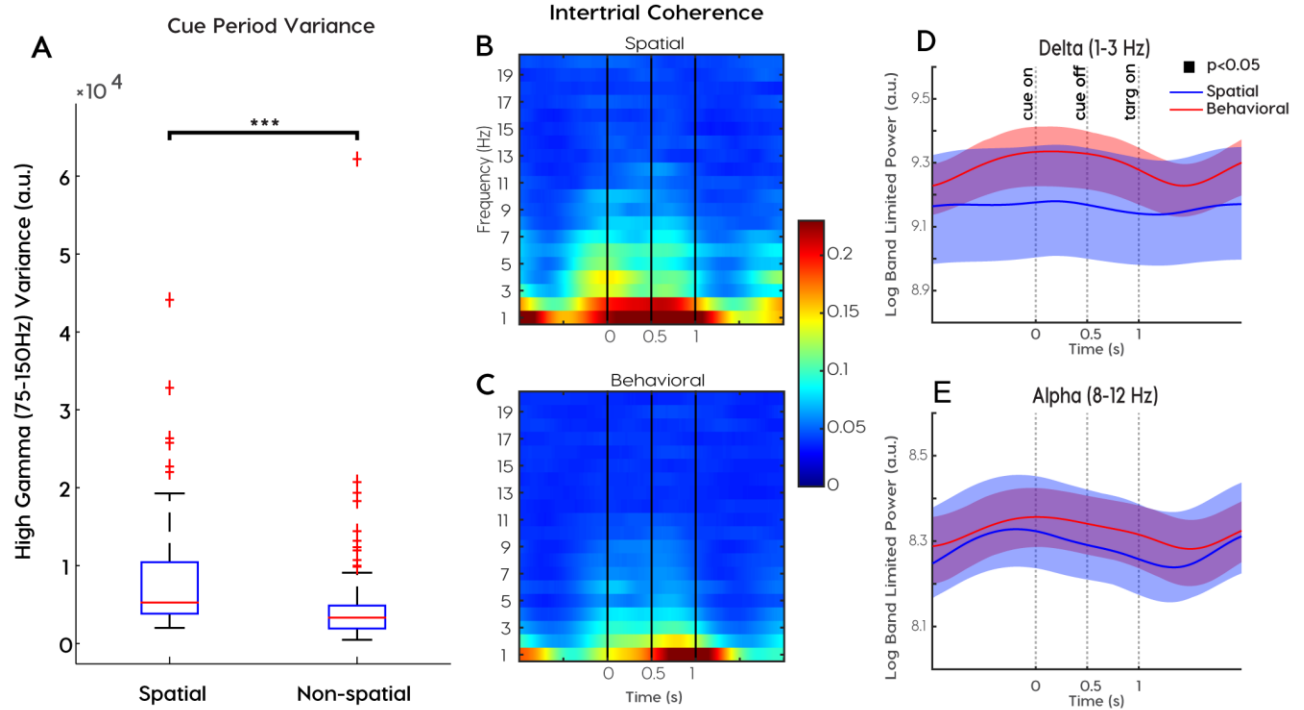
Supplemental Figure 1: Simulating TABL PAC. Simulation of 8 Hz phase coupled with 28 Hz amplitude (red arrows) produces (A) a power spectrum with additional 20 and 36 Hz peaks (i.e. 28 Hz - 8 Hz and 28 Hz + 8 Hz, green arrows) where (B) the amplitude envelope of 28 Hz signal peaks at (D) the 0-degree phase of 8 Hz signal. (C) Phase-amplitude probability density plot (gray dashed line = no coupling) and (E) the comodulogram of the simulated signal.



Supplementary Figure 2: TABL PAC resolution required multiple trials. Using recorded physiological data we calculated modulation indices for different data lengths and averaged the resulting comodulograms after normalizing over frequency pairs. More than 10 seconds of data were required to resolve TABL PAC.



Supplementary Figure 3: Cortical recording site coverage and normalized comodulograms. (Top) All recording sites for 6 subjects were projected onto an averaged brain. Some electrode locations appear removed from the brain surface due to the shrinking that occurs from averaging over brains in three-dimensional space. (Bottom) Averaged comodulograms for each subject.



Supplementary Figure 4: Metrics indicating higher neural modulation at spatial than behavioral sites, despite similar low frequency power. (A) High gamma variance was greater in spatial sites than in behavioral sites. ITC was higher in (B) spatial sites than in (C) behavioral sites. (D) Alpha and (E) delta bands showed no differences between behavioral and spatial sites.

Figure 4.6.1 Supplemental figures

5) Alpha-phase Coupling Distinguishes Mechanistically

Distinct Unconscious States

5.1 Introduction

Coupling of neural oscillations has been shown to correlate with cognition and behavior in multiple species, using multiple modalities^{48,75,81,109,110}. One form of coupling called phase-amplitude coupling (PAC) has been proposed a fundamental mechanism for controlling cortical communication^{83,85}. However, this idea has been challenged by examples of spurious PAC^{87,100} and violations to the mathematical constraints of PAC analyses^{74,101}. In this study, we exploit the cognitive differences between conscious and unconscious states to study the role of PAC. Specifically, we investigate PAC changes caused by slow wave sleep (SWS) and Propofol-induced loss of consciousness (PLOC). The gating by inhibition (GBI) hypothesis is particularly relevant to loss-of-consciousness because it suggests that low-frequencies periodically suppress transmission of neural information⁸³. This hypothesis is relevant because SWS has been shown to suppress cortical transmission⁵⁴ and Propofol causes inhibition through GABA-A agonism¹¹¹. Prior studies have explored changes in PAC during SWS^{57,112–114}, changes in PAC during Propofol anesthesia^{115,116}, and differences in SWS and Propofol anesthesia with scalp EEG⁶². However, no study to date has compared differences in PAC across these two distinct forms of loss-of-consciousness using invasive electrophysiology in humans.

Sleep is organized by the interplay of suprachiasmatic nucleus cells and a complex set of regional networks in the hypothalamus¹¹⁷. It has been shown in human TMS studies that cortical excitability is diminished during sleep⁵⁴. Despite this, sleep is not a passive brain state, but an

active process at both subcortical and cortical levels ¹¹⁸. Accordingly, many electrophysiological features can be measured at the scalp during sleep. These features include slow waves, sleep spindles and k-complexes, each with a well-studied circuitry ^{119,120}. In a recent invasive electrophysiological sleep study, PAC was greater during SWS, when compared to wake, for delta-, theta- and alpha-phases coupled to 30-260 Hz amplitude ¹¹². This result conflicts with evidence that increased PAC facilitates attention ^{48,81}, since attention is markedly reduced during SWS ⁵⁸. More specifically, alpha-phase has a prominent role alertness ^{121,122}, perception ⁴⁷, and selective attention, ¹²³ presumably through an inhibitory mechanism ^{19,106}. These findings are difficult to reconcile with increased alpha-phase PAC during SWS. For these reasons, we hypothesized that alpha-phase PAC scales with inhibitory attention mechanisms and decreases during SWS.

PLOC shares several physiological similarities with sleep. For example, sleep debt, the cumulative requirement for sleep caused by sleep deprivation, facilitates Propofol induction and Propofol anesthesia relieves sleep debt ^{59,60}. Furthermore, slow waves are present in both SWS and PLOC recordings ¹²⁴. Despite these similarities, it is considerably more difficult to arouse someone from PLOC, compared to SWS. Additional electrophysiological differences include the failure of Propofol-induced slow waves to entrain high frequency activity and an anteriorization of alpha activity that occurs in Propofol anesthesia ^{62,64}. PLOC has an inhibitory mechanism of action through GABA-A agonism ^{111,125}. In contrast, evidence suggests that reduced excitability during SWS is due to disfacilitation caused by potassium leak currents rather than GABA-A mediated inhibitory postsynaptic potentials ¹²⁶. Comparing PAC between SWS and PLOC provide a unique window into the role of PAC in the brain.

In this study, we measure PAC in 10 subjects with invasively monitored electrocorticography. Four subjects were monitored during sleep and awake states, five subjects were monitored during anesthetic induction using Propofol and one subject was monitored in both conditions. We hypothesized that alpha-phase coupling indexes inhibitory gating and would therefore discriminate the mechanistic differences between PLOC and SWS. We measure PAC in both conscious and unconscious states using the modulation index described by Tort et al.⁷³. In contrast to prior studies¹¹², we found a double dissociation between sleep state and phase-frequency. Delta-phase coupling increased, but alpha-phase coupling decreased during SWS compared to wakefulness. [We confirmed this finding in an open EEG sleep dataset with 29 subjects.] Importantly, we found that this double dissociation did not occur during PLOC. PAC involving theta- and alpha-phase increased during PLOC compared to wakefulness. We conclude that coupling to alpha-phase distinguishes SWS from PLOC and that this increased alpha-PAC may indicate Propofol-induced cortical inhibition.

5.2 Results

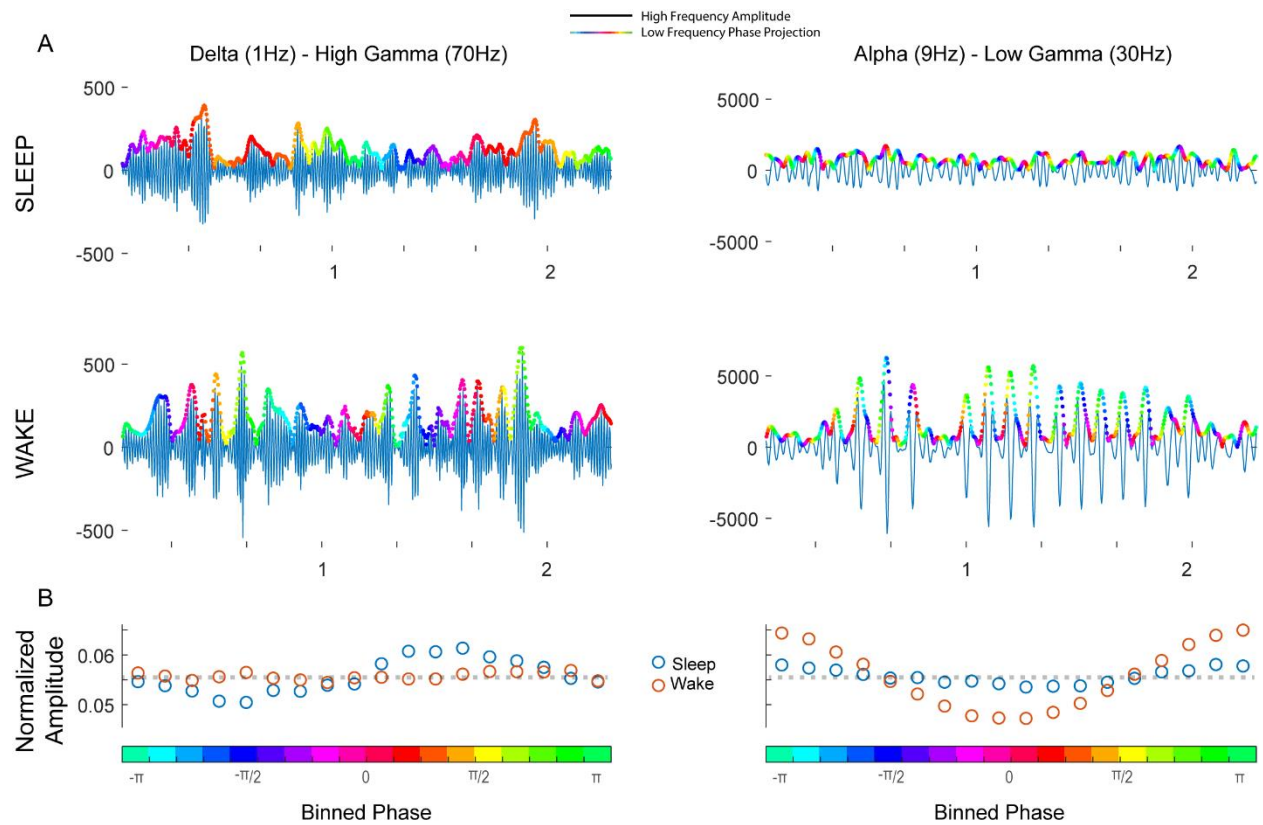


Figure 5.2.1: Double dissociation of PAC by frequency pair and sleep state. (A) Examples of low frequency phase bins (colors) projected on the amplitude envelop of band-limited high frequency amplitude (blue trace) for delta-high gamma (left) and alpha-low gamma (right) in sleep (top) and wake (bottom) states. (B) Normalized amplitude envelopes from all wake (orange) and sleep (blue) recordings binned by low frequency phase (x-axis colors). Grey dashed line represents the uniform distribution or no coupling.

Double dissociation of delta-phase and alpha-phase PAC across wake and sleep states:

We evaluated the difference in PAC between slow wave sleep (SWS) and wake. Figure 5.2.1 demonstrates exemplar coupling patterns elucidating differences between SWS and wakefulness in alpha- and delta-phase PAC. Delta-high gamma (1 Hz – 70 Hz) PAC was greater during sleep (**Figure 5.2.1A top left**) than during wakefulness (**Figure 5.2.1A bottom left**). For visualization purposes, we color-coded the phase of the lower frequency (i.e. 1 Hz) on the amplitude envelope of high-gamma. If the high frequency amplitude was consistently higher on a particular phase of the low frequency signal, coupling existed. The magnitude of delta-high gamma coupling is quantified by the modulation index, or the non-uniformity in the phase-amplitude probability density plot (**Figure 5.2.1e**). Differences in delta-high gamma coupling can be observed between sleep and wakefulness (**Figure 5.2.1A left**). This difference is quantified in the phase-amplitude probability density (**Figure 5.2.1f, red vs blue**).

Alpha-phase PAC had the opposite relationship to SWS. We found an absence of alpha-low-gamma (9Hz – 30Hz) coupling during SWS (**Figure 5.2.1b**) and higher coupling during wakefulness (**Figure 5.2.1d**). This difference is quantified on the phase-amplitude plot (**Figure 5.2.1f, red vs blue**). To evaluate the generality of this double dissociation we tested for differences in wake-SWS comodulograms using a cluster-based permutation method and found significant increases in delta-broadband PAC, along with significant decreases in alpha-broadband PAC, during SWS (both $p < 0.001$. see **Supplementary Figure 5.2.1**).

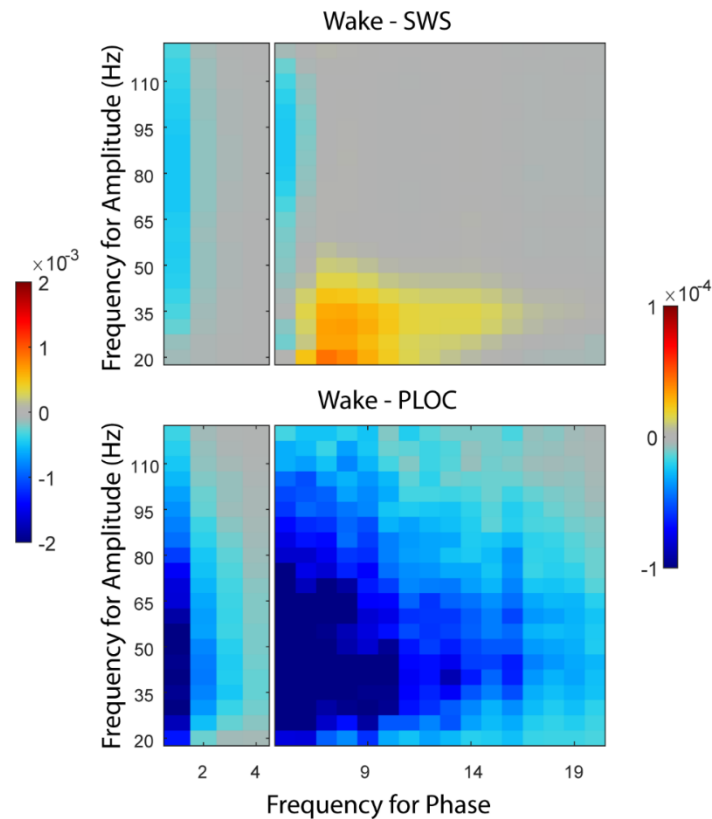


Figure 5.2.2: TABL PAC decreases during SWS but increases during PLOC. (Top) The difference in Wake – SWS comodulograms reveals more TABL PAC in waking state (orange). (Bottom) The difference in Wake – PLOC shows the opposite, a higher TABL PAC in PLOC.

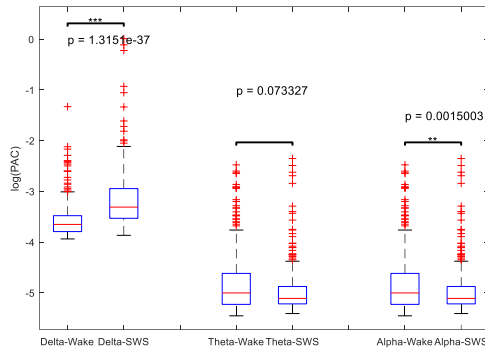
Increase in theta/alpha PAC marks propofol LOC

In stark contrast to SWS, PAC increased dramatically during PLOC and the aggregate change in PAC across cortical sites was consistent. In the transition from wakefulness to SWS there is an increase in delta-phase PAC, no change in theta-phase PAC and a reduction in alpha-phase PAC (**Fig 3A**). In contrast, propofol induction showed a generalized increase in PAC. Delta-, theta- and alpha-phase PAC significantly increased in the last minute of induction, compared to the first (**Fig 3B**). Furthermore, while the increase in delta-phase PAC in both SWS

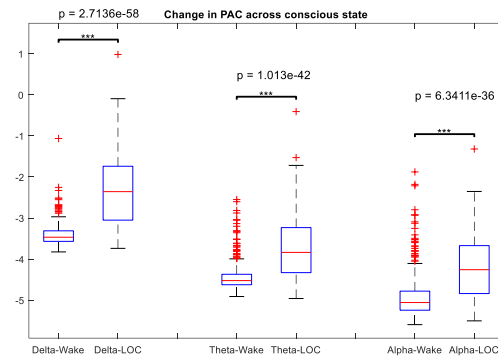
and LOC were accompanied by an increase in power, this was not the case for theta and alpha. Median band limited power using Welch's method demonstrated no change in theta and alpha power in both forms of LOC (**Fig 3 C, D, blue highlight**).

In summary, we have shown that the marked reduction in alpha-phase PAC that occurs in the transition from waking to SWS does not occur in propofol induction. In fact, we see a significant increase in alpha-phase coupling. This increase in coupling cannot be explained by changes in alpha power. Furthermore, we found a larger reduction in beta, low gamma and high gamma in propofol induction, compared to SWS. Therefore, the increase in theta- and alpha-phase coupling is most likely due to the timing of high-frequency bursts of activity (i.e. >25 Hz) relative to the phase of low frequency (i.e. theta and alpha) phase. It is also notable that a) theories on alpha's role in cognition and b) propofol's mechanism of action, both involve cortical inhibition.

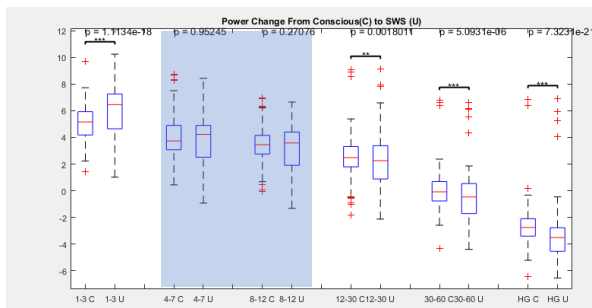
A) SWS Change in PAC



B) Propofol Change in PAC



C) Sleep Change in Power



D) Propofol Change in Power

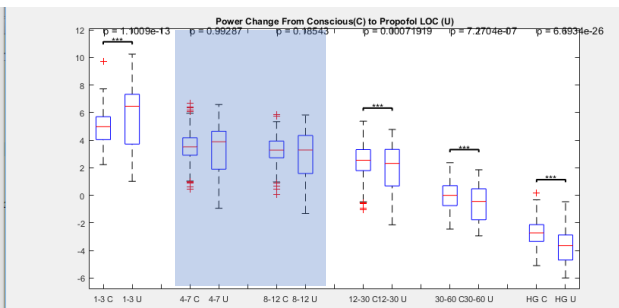


Figure 5.2.3 Increased theta- and alpha-phase PAC is not caused by changes in power. (A) Shows the change in PAC from wake (left boxplots) to SWS (right boxplots). The left two boxes show a positive change in delta-broadband PAC, the middle two boxes display no change in theta-broadband PAC and the right two boxes show a negative change in alpha-broadband PAC. (B) Shows the analogous plot to (A) but for the changes in PAC from wake (left boxplots) to Propofol-induced loss of consciousness (PLOC, right boxplots). We see that delta-, theta- and alpha-broadband PAC increases. (C,D) The corresponding

changes in band-limited power show that delta power increases when transitioning from wakefulness to both SWS and PLOC. Beta, low gamma and high gamma power decrease in both SWS and PLOC, compared to waking. However, theta and alpha power does not change (blue shading).

5.3 Discussion

In this study we investigated whether PAC increases or decreases in two states of reduced arousal SWS and PLOC. First, we identified a double dissociation in PAC frequency pairs and sleep state. During SWS delta-broadband PAC increased but alpha-broadband PAC decreased. In contrast, alpha-broadband PAC increased in PLOC, compared to wake. Therefore, the similarities between SWS and PLOC may be indexed by delta-broadband PAC increases. These include the ability of both SWS and PLOC to relieve sleep debt. The differences between SWS and PLOC may be represented in alpha-broadband PAC. The major difference is the mechanism behind both states. While SWS is related to disfacilitation, PLOC is caused by inhibition.

The differences in the results of our study and the most comparable work in the literature¹²⁷ warrants a discussion of methods. In a personal communication with Amiri et. al., we learned that the authors did not adhere to the mathematical constraints of PAC analyses laid out in prior theoretical literature^{74,128}. Namely, frequency-for-amplitude filters must have twice the bandwidth of the frequency-for-phase of interest. If we want to assess coupling to 8 Hz phase, our amplitude filters must have a 16 Hz bandwidth. Given that Aru et. al. used filters with insufficient bandwidth, it is likely that their results only reflected delta-broadband coupling. Like Amiri et. al., our study also demonstrated increased delta-broadband PAC during in SWS. Our

additional finding regarding PLOC adds to the understanding of how consciousness and PAC are related.

Delta-broadband PAC likely represents the slow waves that are common to both PLOC and SWS. These slow waves may function to periodically disfacilitate neuronal firing. This type of globally disfacilitated activity may be a marker of low arousal states. On the other hand, alpha-broadband PAC may specifically represent inhibition. Prior studies have established that SWS is not an inhibitory state¹²⁶. In contrast, PLOC has a clear inhibitory mechanism of action, GABA-A agonism. Taken together, delta-broadband and alpha-broadband PAC can discriminate waking, SWS and PLOC states along the axes of arousal and inhibition. This new correlate of conscious state may appear perplexing. Why should waking be more similar to PLOC than SWS along any axis when behaviorally PLOC and SWS are quite similar? While this topic is beyond the scope of this research we offer one clinical paradox that may be elucidated by this finding.

Little is known about recovering consciousness in patients that are in vegetative or minimally conscious states. However, evidence exists that Zolpidem (i.e. Ambien), a GABA-A agonist like Propofol, can transiently recover consciousness in these patients⁵⁰. Prior research in attention suggested that TABL PAC correlates with inhibition (see Chapter 4). Considering the plethora of information our brains process every moment, we propose the following hypothesis for why GABA-A agonists may clinically improve consciousness. An inability to sift through the plethora of information our brain receives may result in an inability to engage in goal-directed behaviors. Inhibition through GABA-A agonism may temper the onslaught of information just enough for productive trains-of-thought to be selected and acted upon. Non-invasive measurements of TABL PAC before and after Zolpidem treatment in vegetative patients may quantify this change in future experiments.

5.4 Supplemental Data

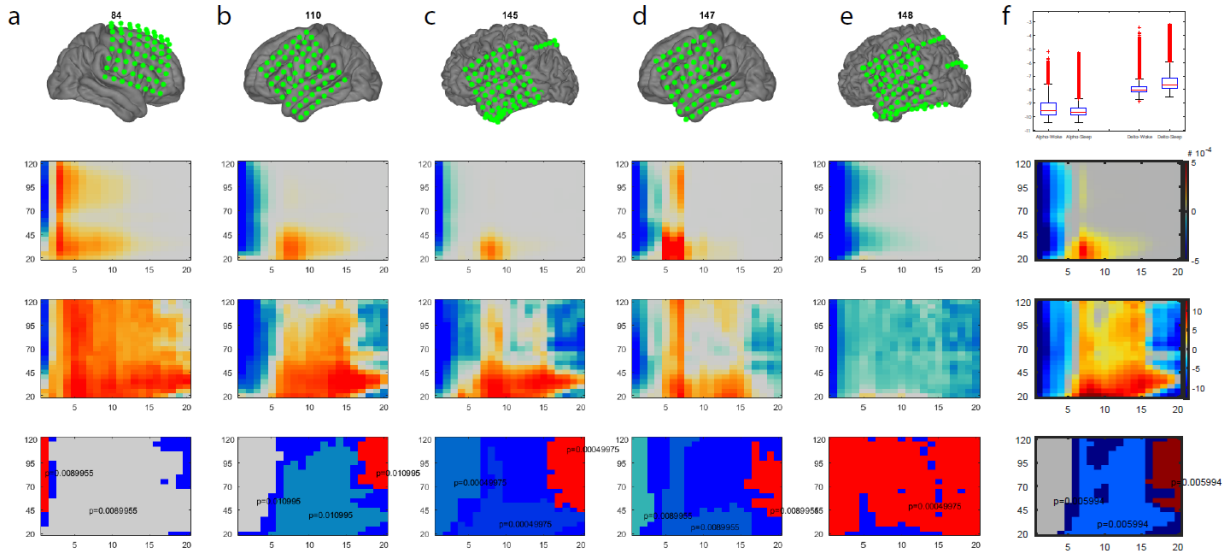


Figure 5.4.1: Consistency of double dissociation across patients. (Top) ECoG grid placements for all subjects. (2nd Row) Wake - Sleep Comodulograms. (3rd row) T-statistics for Wake – Sleep across all recording sites. (Bottom Row) Cluster-based permutation testing reveals significant clusters (red, grey, teal and blue). Frequency-for-amplitude on y-axis and frequency-for-phase on the x-axis for bottom 3 rows.

Consistent dissociation by phase-frequency and sleep state across subjects:

Comodulograms represent the difference in coupling (i.e. non-uniformity of phase-amplitude plots), across waking and SWS states (i.e. wake – sleep), for every pair of frequencies under investigation. We averaged these comodulograms across all cortical sites both within and across subjects. The five subjects providing sleep and wake data had ECoG grids placed over or near the Sylvian fissure, with coverage over parts of the temporal, pre-motor, motor, and sensory

cortices. ECoG strips were also placed over parieto-occipital regions and ventral regions (**Supplemental Figure 5.4.1a-e, top row**). To compare SWS and wake states, we observed the difference in comodulograms (**Supplemental Figure 5.4.1a-e, middle top row**), the t-statistics (**Supplemental Figure 5.4.1a-e, middle bottom row**) and the cluster-based statistical inference (**Supplemental Figure 5.4.1a-e, bottom row**) for each subject⁹². All subjects displayed an increase in delta-phase PAC in SWS compared to wake. Conversely, alpha-broadband PAC was decreased in SWS, compared to wake. The t-statistic tested the hypothesis that there was no difference between wake and SWS states and uses the pooled standard error across wake and SWS for every frequency-pair in the comodulogram. The t-statistics partly rectify the 1/f power-law distribution and were more sensitive to small, yet stable, differences in PAC at higher phase frequencies (**Supplemental Figure 5.4.1a-e, middle bottom row**). When we combined the data from all subjects, we observed the same dissociation between delta-broadband and alpha-broadband PAC across wake and SWS (**Supplemental Figure 5.4.1f, middle rows**).

We used a cluster-based permutation test to assess the significance of these differences. Adjacent t-statistics above or below a two-tailed threshold ($\alpha = 97.5$) were summed to generate cluster-statistics. Cluster-statistics were tested against a permutation distribution. Permutation distributions were created by shuffling wake and SWS labels 1000 times and repeating the procedure for each shuffle. Only the maximum cluster-statistic was used in the creation of the permutation distribution. In four of five subjects, a significant cluster associated with delta-phase emerged distinctly from a significant cluster associated with alpha-phase. The border between these clusters ranged from 2-7 Hz and both clusters included the entirety of the frequency-for-amplitude spectrum (20-150 Hz). In the fifth subject (i.e. 148), there was no alpha-phase cluster. Only one cluster was found that ranged the entire phase and amplitude spectrum.

However, this subject had a notable region of non-significance in the theta-alpha-phase (5-15 Hz), beta-low-gamma-amplitude (20-45 Hz) region of the comodulogram. All clusters passed a threshold of $p < 0.05$, correcting for comparisons using the false discovery rate (FDR) correction for multiple comparisons (**Supplemental Figure 5.4.1a-e bottom row**).

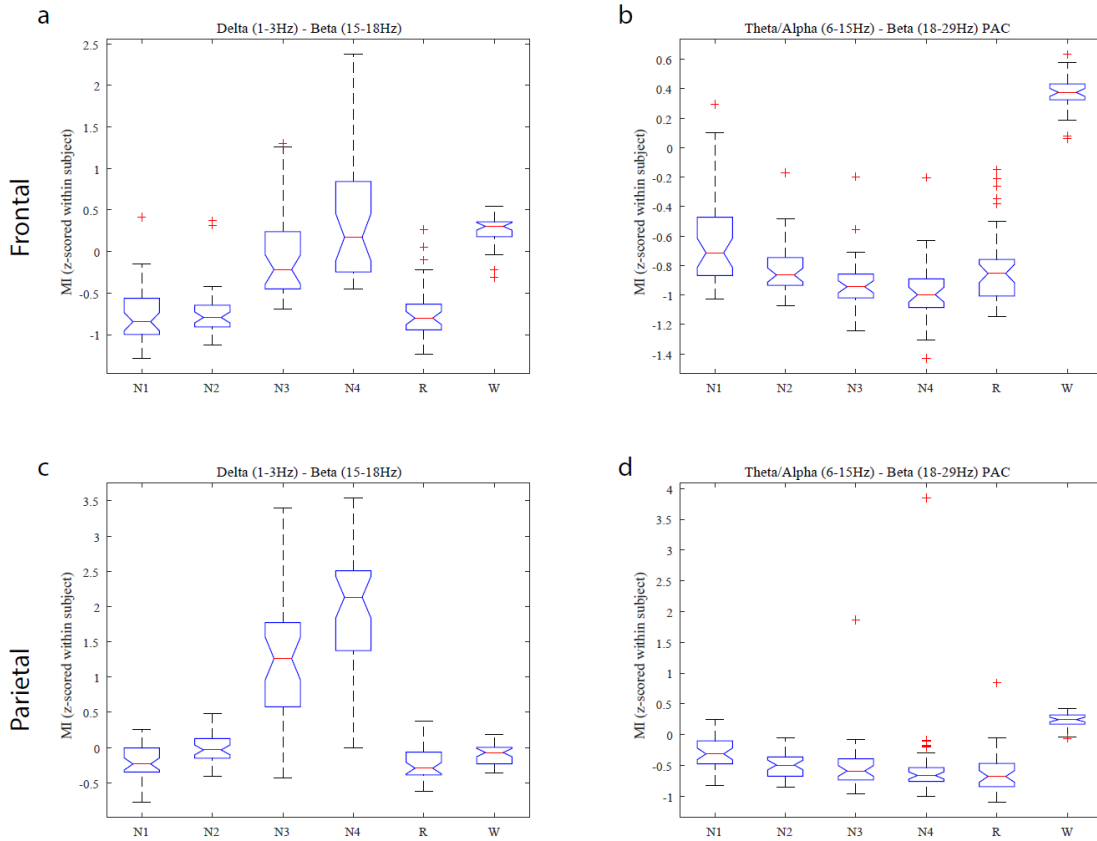
We pooled the data from all participants to perform a group-level statistical inference using the cluster-based permutation test mentioned above (**Supplemental Figure 5.4.1f, bottom row**). Here we again found a division between delta-phase and alpha-phase significant clusters. We used the delta-phase and alpha-phase clusters as spectral regions-of-interest (ROIs) and summed the modulation indices within these spectral ROIs for every cortical site in every participant. Plotting these delta- and alpha-phase PAC distributions for sleep and wake states demonstrates the generality of the reversal shown in Figure 5.2.1. Delta-phase PAC is significantly higher during SWS, compared to wake (rank sum, z -value = 7.2, $p = 5e-3$) and alpha-phase PAC is significantly lower during SWS, compared to wake (rank sum, z -value = -30.6, $p = 1.8e-205$) (**Supplemental Figure 5.4.1f, top row**). In summary, we found a consistent reversal of delta-phase and alpha-phase PAC across SWS and wake states.

EEG scalp recordings and sleep staging

One drawback to our ECoG analysis was the lack of sleep staging, which is commonly done with scalp EEG recordings during sleep studies. To understand PAC dynamics across sleep stage, we employed the dataset posted by Kemp et. al. on the EDF database hosted by PhysioBank^{129,130}. Despite known differences in spectral range and spatial specificity across scalp EEG and intracranial ECoG, we hypothesized a reversal in delta- and alpha-phase PAC

phenomena between wake and sleep states. The Kemp et. al. dataset was recorded at 100Hz and includes two leads, both are centrally located along the lateral axis while one was located frontally and the other parieto-occipally.

We plotted the average change in modulation index for the frequencies of interest using all 29 subjects across sleep state. We observed several similarities to the ECoG data. Specifically, we found an increase in delta (1-3 Hz)-broadband PAC during N4 sleep, for amplitudes less than 20 Hz. This sleep-related delta-phase PAC was observed most strongly in parieto-occipital sites during N4 sleep. Conversely, we found a theta-alpha-phase PAC signal that steadily decreased across N1-N4 sleep stages in parieto-occipital sites. This theta-alpha-phase PAC varied in intensity across phase-frequencies for frontal and parieto-occipital sites. In parieto-occipital sites, however, we found a reduction in alpha-phase PAC in the N2 stage of sleep which remained through every subsequent sleep stage, including REM sleep (i.e. N3, N4 and R). We illustrated this reversal of delta- and alpha-phase PAC by plotting respective PAC distributions for both frontal and parietal sites (**Supplementary Figure 2**).



Supplementary Figure 5.4.2: Delta-phase PAC increases, and alpha-phase PAC decreases, with increasing NREM sleep stage. We performed our ECoG analysis on EEG data that was recorded by another group of researchers. In the parietal EEG electrode we found the same relationship, an increase in delta-broadband PAC during SWS and a decreased in alpha-broadband PAC. This generalizes our ECoG results to non-invasive electrophysiology.

In summary, we demonstrated that the delta-/alpha-phase PAC reversal found in ECoG between SWS and wake states was also observed in EEG data collected by separate group of researchers. Differences between EEG and ECoG, (including signal leakage and smearing plus the resistance of skull and scalp that results in an inability to record high gamma signals from the scalp with EEG), could have caused divergence in the dataset such as muscle and motion artifact.

However, this finding is important because recent reports have suggested that theta- and alpha-phase PAC increases across sleep stages N1-N3¹¹². We did not find this to be the case. From sleep stages N1 to N3, delta-phase/low beta-amplitude PAC monotonically increased, but alpha-phase/high-beta amplitude PAC monotonically decreased.

6) Proof of Concept and Conclusion

The goal of this project was to develop control features for a hemineglect BCI. We began by realizing that, unlike the motor system, attention is a multifaceted cognitive function that is distributed in the brain. In the previous two chapters we characterized electrophysiological features representing a) lateralized covert attention, b) reaction time and c) arousal, each of which are problematic for the hemineglect patient. Human brains are highly variable, so attentional control features may differ in frequency and spatial location from person to person. A next step in developing a hemineglect BCI is to use a screening task to determine and optimize control features for each patient. After these features are identified, they can be used in real-time feedback. Theoretically, a patient will learn to control these features and thereby improve their attentional deficit. In prior work, Gunduz et.al. classified the locus of attention with ECoG in one of three positions with above-chance accuracy¹³¹. Highly weighted features included alpha, beta and gamma activity over posterior parietal cortex, premotor cortex, dorsolateral prefrontal cortex and frontal eye fields^{131,132}. Here we extend prior work by predicting different aspects of attention in a post-hoc cross-validated machine learning analysis. We will analyze the Posner task and the sleep dataset as if it were a screening task and see how well we can predict attention and arousal on a single trial basis.

Predicting Spatial Attention: Figure 6.1.1 is proof of concept for predicting lateralized covert spatial attention, which is most noticeably deficient in hemineglect. Using trials from our Posner task (see Chapter 4) we developed a log-loss classifier to predict when a subject was covertly attending to the left or right. We divide all the trials into a testing and training dataset. We train our classifier on the training dataset and used the resultant weightings to predict labels

in the testing dataset. We do this 10 times, with 10% of the data used for testing, to cross-validate the accuracy of our algorithm. After minimal parameter optimization, the accuracy of this algorithm was high (73%) and significantly greater than chance. Not all subjects yielded classification accuracies that were above chance, which may reflect differences in placement of the ECoG grid and individual variations in recording quality or behavioral performance. Previous attempts at classifying the locus of attention using EEG made use of the difference in activity across hemispheres⁴¹. ECoG is rarely collected from both hemispheres simultaneously but future EEG approaches may include this difference across hemispheres as a control feature. Furthermore, these classification attempts did not make use of PAC or sharp waves. Future attempts might add these novel features to the classifier and test if performance increases.

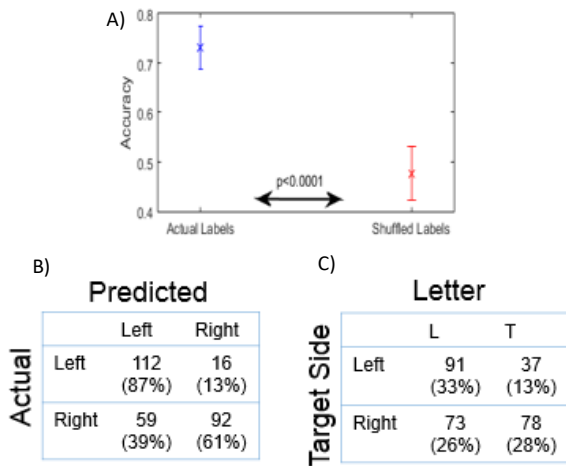


Figure 6.1.1: Log-loss classifier performance for L vs R target locations. Correct trials with the highest 90% of reaction times were used for classification. The dimensionality of ECoG signal during target presentation was reduced with PCA and a log-loss classifier was trained on target location (Left = -1, Right = 1). The results of ten-fold cross validation are shown. The classifier had 73% accuracy at classifying the location of targets (A). Shuffled labels were used as a control and confirmed chance classifier estimates. The classifier performed worse on right trials (B). A slight correlation between target letter and side was observed for left sided trials (C) due to selection criteria.

Predicting Reaction Time: We predicted RTs using sharp waves and band-limited theta, beta and alpha power from the cue period as features. We utilized a ridge regression algorithm with L1 regularization in a leave-one-out cross-validation procedure. We used principle component analysis to reduce the dimensionality while accounting for 90% of the variance in the dataset.

Using only behavioral sites yielded 68% accuracy in predicting whether a trial would be in the top or bottom 3rd of RTs. Using all cortical sites yielded 69% accuracy, while spatial sites only yielded 63% accuracy. Our reduced accuracy at predicting fast vs slow trials may reflect the multiple factors reflected in RT. A host of factors beyond attention and arousal may influence reaction time such as vigilance, motor preparedness, improved object identification, visual acuity and so on. Future work may benefit from considering the relationship between RT and cognitive functions beyond attention.

Predicting Arousal: The sleep dataset from Chapter 5 included ECoG recordings from 5 patients (with roughly 64 channels each) with several hundreds of minutes in sleep and wake data. Using only the comodulograms generated on single minutes of data, we divided the dataset into testing and training sets and classified sleep using a support vector machine. This procedure yielded near-perfect accuracy in predicting sleep vs wake conditions. For all subjects we predicted sleep vs awake with 99+% accuracy. This robust classification accuracy was maintained even when we trained with 1% of the data available. Future work should compare the accuracy of PAC classifications to standard sleep staging algorithms.

Taken together, these data complete the first steps toward a hemineglect BCI, but questions remain. We established that the Posner task can be used to detect cortical regions related to both the spatial and non-spatial aspect of the hemineglect deficit. Furthermore, these results suggest that PAC can be used to detect slow wave sleep. Future work might focus on parameter optimization and comparing the accuracy of different algorithms. As future work transitions from researching attention in healthy patients to focusing on hemineglect patients, several questions will become important. Firstly, how do attentional control signals change in hemineglect? We learned from motor BCIs that if a patient damages her right motor cortex, her

left motor cortex can compensate. It is unclear if there are compensatory mechanisms in hemineglect that can be used in the same way. Secondly, how do attentional control signals interact? Evidence suggests that drowsiness or arousal interacts with spatial attention. When healthy people are falling asleep they experience a shift in their spatial bias¹³³. Understanding how control signals interact will be important to optimize feedback. Finally, it remains unclear how multiple control signals can be used in the context of BCI neurofeedback. Typically one control feature is used for training. For example, the IpsiHand BCI uses a control feature for hand-opening. It is unclear whether control features can be used for BCI training independently, or if they will have to somehow be combined into a single control feature. Answering these questions will be important in the development of BCI neurorehabilitation for hemineglect.

Conclusions: In this work we set out to identify electrophysiological correlates of attention that could be used in a hemineglect BCI. We found a novel form of PAC, theta-alpha-phase coupled to beta-low gamma-amplitude (TABL), that correlated with RT. TABL PAC was related to transient beta (50 ms) waves that occurred at roughly 140 ms intervals. We developed a method to identify these transient waves in short periods of time. Furthermore, TABL PAC also distinguished between different states of arousal such as wakefulness, slow wave sleep and Propofol anesthesia. Our clinical and engineering goal was to move towards a hemineglect BCI. As a proof-of-concept we used band-limited power, sharp waves and TABL PAC to predict spatial attention, reaction time and sleep state with above-chance accuracy. Our scientific goal was to better understand how attention was represented in the brain and, more generally, to understand the relationship between attention and other states of consciousness. We found that TABL PAC was higher during wakefulness than sleep, but also found high TABL PAC during

Propofol anesthesia. This leads line of evidence suggests a common link between high and low arousal states, inhibition.

In identifying how different forms of attention were encoded in the brain, we concluded that TABL PAC was inhibitory for several reasons. First, alpha and theta have previously been shown to be involved in inhibition^{134,135}. Second, the 180-degree phase preference of TABL PAC is just another way to describe “pulsed inhibition”^{47,106}. Third, TABL PAC was greater in sites with less neural modulation, as indexed by ITC and high gamma variability. Interestingly, we did not mention in Chapter 4 that TABL PAC was highest in the sites that were neither spatially or behaviorally relevant. Fourth, TABL PAC magnitude was highest in sites with relatively less broadband power. Fifth, TABL PAC was high during Propofol anesthesia, a GABA-A inhibitory agonist, but was low during sleep. This may seem surprising at first, but slow wave sleep is a disfacilitated, rather than inhibited, state. Putting all this together allows us to append our working model of consciousness. Previously we spoke the “spectrum of consciousness” with a two-dimensional axis: awareness and arousal. Propofol-induced loss of consciousness is at the bottom left, where arousal and awareness are lowest (**Figure 6.2 red**). Sleep is higher and to the right (**Figure 6.2 blue**), because people are more arousable during sleep and can respond to stimuli. Finally, attentional states are at the very top right of the plot (**Figure 6.2 green**). Considering our TABL PAC results we can add a third dimension: inhibition. This third dimension shows us that the spectrum curves so that attentional states are closer to Propofol-induced loss of consciousness than sleep along the “inhibition” access. This has been predicted in previous theories that suggested the first step in attending is globally suppressing neural activity. Only after global inhibition can relevant cortical areas can be selectively disinhibited⁸⁸. Finally, the addition of the inhibition dimension explains how patients in

vegetative states can be re-awakened with GABA-A agonists like Ambien. If too little inhibition occurs, then a patient may be in a sleep-like state. Too much inhibition and the patient is knocked out.

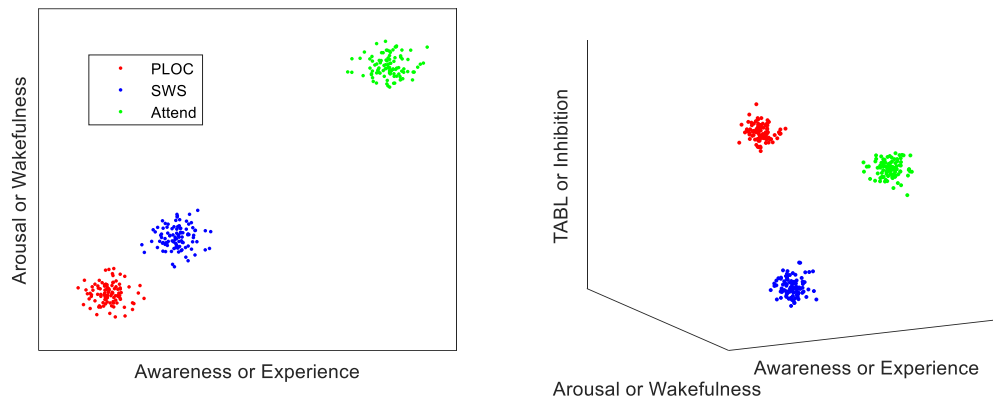


Figure 6.1.2 *Adding a dimension to the spectrum of consciousness. Here we propose that TABL PAC is a marker of inhibition. TABL PAC is high in waking (green) and Propofol (red) states, but not during SWS (blue).*

In conclusion, this work takes several important steps in our scientific understanding of attention and in the practice of building BCIs to augment it. The continuation of this work is promising. In the future, we will send patients home with EEG devices that will automatically determine correlates of attention, arousal, and, possibly, consciousness. This will benefit hemineglect patients, but it may also help healthy individuals learn and complete tasks. Ultimately, this may one day realize Edward Mitchell’s vision of imbuing “superhuman power(s)” with brain computer interfaces.

Appendix A: A Review of Rehabilitation Paradigms for Hemineglect

Rehabilitation techniques for HN have been divided into bottom-up and top-down approaches. Bottom-up techniques include cold-water calorics, optokinetic stimulation and prism goggles. Bottom-up techniques target primary sensory areas and reflexes. They are practical because they don't require the patient's awareness to work. HN patients are often unaware of their behavior (anosognosia). Top-down approaches, on the other hand, require deficit awareness and encourage voluntary compensatory strategies. These include, cueing, visual scanning or mental imagery¹⁴. We will briefly review these approaches.

Cold-water Calorics and Optokinetic Stimulation: Introducing cold water into the left ear of HN patients elicits a vestibulo-ocular reflex causing nystagmus towards the irrigated ear. This temporarily improves gaze deviation in HN. Additionally, this intervention improved other non-visual deficits like tactile hemianesthesia and anosognosia for left hemiplegia¹³⁶. It is notable that this form of bottom-up stimulation improves non-spatial deficits like anosognosia. A similar mechanism is thought to underlie optokinetic stimulation. This paradigm consists of many dots randomly moving right to left on a screen. Dot speeds were designed to generate the greatest amount of nystagmus¹³⁷.

Prism-Adaptation: The most successful bottom-up rehabilitation technique is prism adaptation. Prism goggles are worn that bend the light path incident on the eye toward the wearer's right. The goggles cause wearers to initially overshoot to the left when pointing at objects in front of them. This scene that is incident on the wearer's eyes is to the right of where they are facing. Adaptation occurs when the wearer learns this new visuo-motor transformation and pointing accuracy improves. After adaptation, the goggles are removed resulting in a rebound leftward bias in pointing movements. This bias is toward the neglected hemifield of HN patients. After adaptation training HN patients reported more contralesional stimuli¹³⁸. A two week rehabilitation paradigm showed effects up to 5 weeks post treatment¹⁵. Interestingly, HN patients can only adapt to prism goggles that are bent right and not to those bent left.

Cueing and Feedback: Therapists commonly cue HN patients to attend to contralesional space during daily activities like eating, dressing and cleaning. Caregivers of HN patients naturally cue them to important things they've neglected such as food on their plate. It is also used in a number of rehabilitation paradigms. These involve a) strategies of placing and actively locating the contralesional arm and b) stimulating that arm periodically on a variable time interval requiring response¹³⁹. These interventions reduced HN symptoms in standard neglect tests. One study used eye-tracking glasses to deliver auditory feedback on eye-movements¹⁴⁰. A different tone was played for leftward and rightward eye movements. This device was both enjoyable and effective

in the short term. In 1994, some authors argued that the size of the device limits frequent use¹⁴¹. The ubiquity of portable electronics in recent years points to more opportunities for automated cueing or feedback devices. This is especially true for HN patients who have anosognosia, a lack of awareness about their deficit.

Visual Scanning: Neglect patients show disordered visual search strategies, reduced reading speeds and bias eye-movements ipsilesionally. Visual scanning a rehabilitation technique that is often combined with feedback on performance. Subjects are required to direct a saccadic eye movement to targets that appear across a screen. Initially targets appear in a predictable fashion. As training progresses, randomness of subsequent target location is introduced making the task more difficult. Subjects are asked to report letters or digits at the target location¹⁴.

Mental Imagery: When HN patients are asked to recall a scene from memory they omit the contralesional elements of that scene. However, after imagining themselves turning around, the HN patient will be able to recall the omitted elements^{16,142}. This indicates that HN symptoms extends to the imagination. “Imagery training” was shown to ameliorate HN performance deficits. Subjects were instructed to describe the contents of familiar rooms, pathways and geographic locations. They were also asked to imagine movements. Although the sample size was small, these interventions showed a positive effect¹⁴³.

Virtual Reality: Castiello et. al. created a virtual reality paradigm for HN where virtual objects occupied one of three locations on a screen. The task was performed on a screen with a virtual arm. Subjects were instructed to move their own arm to one of three positions, identical to the virtual environment. On different trials the virtual arm appeared in locations both congruent and incongruent with the subject’s real arm. An incongruent trial required participants to adjust their strategy and reach to an unoccupied location. The authors hypothesize that a form of visuomotor adaptation occurs in this paradigm that is similar to prism adaptation. The subjects perceived errors in their visuomotor model of the environment, which prompts a remodeling of that environment. This process improves HN symptoms¹⁴⁴.

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