Evidence for Information Processing in the Brain
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ABSTRACT

Many cognitive scientists, neuroscientists, and philosophers of science consider it uncontroversial that the brain processes information. In this work we broadly consider the types of experimental evidence that would support this claim, and find that although physical features of specific brain areas selectively covary with external stimuli or abilities, there is no direct evidence supporting an information processing function of any particular brain area.

1. INTRODUCTION

Many cognitive scientists believe that the brain processes information. Bechtel and Richardson (2010), as philosophers of cognitive science, consider it uncontroversial that cognitive scientists involved in neuroimaging research believe that “the brain contains some regions that are specialized for processing specific types of information (p241).” Neuroscientists too claim that “the principle function of the central nervous system is to represent and transform information (deCharms and Zador 2000, p613).” Given such wide-spread acceptance of a belief, it is appropriate to ask for the justification of this belief. If the justification is empirical and experimental, then we should look to the research reported by working scientists in the field; if it is metaphysical, then we should look to the arguments of philosophers and theoreticians.

We will no doubt discover both kinds of justification if we look for it. Yet we assume that cognitive scientists, when stating that the brain processes information, are primarily stating an empirical fact or a widely agreed-upon scientific proposition that is supported by a body of experimental evidence. Like the physicist who can back up the proposition “protons have spin” with a presentation of the experimental evidence, we expect that the cognitive scientist should be able to do the same regarding a statement about the brain. If the cognitive scientist cannot do this, then the proposition is non-empirical or unscientific. We are not suggesting a definition of science or solving Popper’s demarcation problem, but we are appealing to the belief that accepted scientific statements are associated with experimental evidence. Without associated evidence, a proposition cannot be scientific.

Our task here, however, is somewhat more involved than an objective review of the scientific literature. As we have learned from philosophers of science over the past century, “theory dominates the experimental work from its initial planning up to the finishing touches in the laboratory (Popper 1959, p90).” Hanson (1958) and Kuhn (1962) were among the first to direct our attention to the theory-ladenness of scientific observation. Brewster (2001) extended this position, arguing that the complete scientific process, which includes attention, perception, data interpretation, memory, and scientific communication, is influenced by theory. Perhaps most relevant to our work is Popper’s warning that “…observation statements and statements of experimental results, are always interpretations of the facts observed…they are interpretations in the light of theories (italics in original, p. 90).” Of course, none of this need imply scientific relativism, and relativism is not assumed in this work.

One may presume that only cognitive scientists are qualified to interpret the experimental evidence in the field. While an expert’s assessment carries more weight than the non-involved observer, it is reasonable that anyone who takes time to understand the evidence and its methods of acquisition is in a position to construct an interpretation. The force of the interpretation should be based upon the reason of the argument and not only its source. Nonetheless, we have performed some of the types of experiments that we are now interpreting.

Cognitive scientific evidence, especially neuroimaging evidence, has been increasingly subjected to criticisms. To better demarcate our position, we highlight that we are not specifically arguing between distributed versus localized processing in the brain (Utel 2001, Hardcastle and Stewart 2002), and we are not pointing out the previously discussed technical-methodological limitations of brain assessing technologies (Logothetis 2008, Roskies 2007, Klein 2009). We do share with these authors the broader concern for interpretations of evidence in the field of cognitive science, and how theoretical assumptions influence interpretations of evidence, ultimately ending in statements made by cognitive scientists that carry the weight of scientific fact. These facts, in turn, are used by naturalistic philosophers of mind to constrain philosophical theory and argument.

2. COGNITIVE SCIENCE EVIDENCE TYPES

The scientific statement that we will consider is Bechtel and Richardson’s proposition “the brain contains some regions that are specialized for processing specific types of information,” although, since we are not specifically arguing against localization of brain function, we will consider simultaneously the more general proposition P “the brain processes
information.” deCharms and Zador say that it is the function of the brain to process (represent and transform) information. There is no philosophical consensus on how to define a natural biological function, and we will assume that processing information is a natural function of the brain like pumping blood is a natural function of the heart. One might argue that processing information is a natural process and not a natural function, and we will consider both alternatives, at times blurring the distinction in this work.

We wish to consider the experimental evidence that justifies P. There are tens of thousands of papers in scientific journals that may be used as evidence, thus a systematic evaluation of every paper independently and subsequent integration of the evidence is not feasible. The task must be simplified, but in a way that addresses the initial question. As a first step, we will only consider research that involves measuring or manipulating the physical properties of brain. While a study that does not involve brain properties may contribute to our scientific understanding of the brain, it can only do so indirectly by prompting theory formation and characterizing behavioral phenomena. For example, in 1908 Yerkes and Dodson discovered that performance on a task at first increases with increasing arousal and then decreases once arousal levels become too great. They quantified the intuition that ‘stress’ can enhance performance. This interesting and useful finding may suggest neurophysiological correlates of performance and arousal, but it does not experimentally justify any statement about brain function in a direct sense.

We are primarily interested with research that investigates the relations between physical brain properties and behaviors, abilities, or physical (sensory) contexts. Cognitive scientists, and philosophers of science, reference the evidence from this category of research when making claims about brain function. Relational evidence, as we will call it, can be broken down into four major categories:

1) **structure/ability studies**,
2) **external-stimulus/brain-response studies**,
3) **task/brain-response studies**, and
4) **brain-manipulation/behavioral-response studies**.

In structure/ability (or S/A) studies, researchers relate the structure or structural states of the brain to the absence or presence of particular behaviors or abilities. Paul Broca (1861) popularized this type of research with his lesion-deficit, or lesion study, when he discovered an individual who could only speak the syllable ‘tan.’ A post-mortem analysis of the person’s brain revealed damaged brain tissue in the posterior part of the left inferior frontal gyrus—a region now known as Broca’s area. Thus Broca related the ability to produce fluent speech to the left inferior frontal gyrus. The class of S/A studies includes more than lesion studies since any physical feature of the brain (e.g. patterns of white matter connectivity) may be associated with the absence or presence of specific abilities.

The structure in S/A studies refers to the physical structure of the brain as measured by a variety of measuring techniques, the most basic being gross anatomical observation of brain tissue. Other measuring techniques include, but are not limited to, histological examination, molecular analysis, electroencephalography (EEG), magnetoencephalography (MEG), computed tomography (CT), positron emissions tomography (PET), single photon emitted computed tomography (SPECT), structural magnetic resonance imaging (MRI), diffusion tensor imaging (DTI), magnetic resonance spectroscopy (MRS), and others. By ability we mean any observable behavior that can be done by an animal or human, such as the ability to count out loud, to raise one’s arm, to navigate a maze, to write a sentence, to score above chance on a test, etc.

In external-stimulus/brain-response (ES/BR) studies, the experimenter systematically manipulates a physical feature of an organism’s external environment, and measures temporally coincident properties of the organism’s brain. The response need not occur precisely simultaneous with the stimulus and is typically extended in time. Edgar Adrian is generally credited with pioneering stimulus-response studies of nervous tissue. He was the first to record the electrical activity of single nerve fibers, and was subsequently awarded a Nobel Prize in 1932 for his work. As an example of his prolific research, he isolated an eel’s eye and optic nerve, attached electrodes to the nerve, and recorded the electrical activity to varying lighting situations (Adrian and Matthews 1927).

Brain responses in ES/BR studies are recorded using a variety of techniques based upon electromagnetic brain properties, including single-unit intra and extracellular recording, evoked potentials, EEG, MEG, and others. Functional MRI (fMRI) is a popular tool used by cognitive scientists to assess brain properties in response to an ES. Proper interpretation of the fMRI signal itself requires technical background knowledge (Roskies 2007, Logothetis 2008); briefly, the fMRI signal is consequence of the magnetic properties of blood components which covary with local metabolic demands. Other common techniques in ES/BR studies include PET and optical imaging.

When, within a research protocol, a brain-response is recorded while the organism is performing a particular task or activity, we call this a task/brain-response (T/BR) study. The form of the T/BR study is similar to ES/BR studies, except the process of completing the task is ‘self-directed’ rather than under complete control of the experimenter. Often T/BR studies include ES aspects as well. Memory research provides typical examples. Poppenk et al. (2010), in studying prospective memory which is described as the ability to act out postponed intentions at future times, presented a series of visual scenes (ES) to subjects and instructed the subjects to either imagine performing an action associated with the visual scene, or to use the scene as a reminder to perform an action the next time the same scene was viewed. FMRI was used to measure properties of the subjects’ brains during the tasks. Notice that although the experimenter controls the ES and the task command directly, she cannot control the process by which the subject completes the task.

Brain-manipulation/behavioral response (BM/BR) studies differ from ES/BR and T/BR studies in that physical properties of the brain are directly controlled or manipulated while a behavioral response is observed. Technologies used for BM
include lesioning, gene-expression, direct current stimulation, 
electrode-based deep brain stimulation, transcranial magnetic 
stimulation, and light-based optogenetic stimulation, among 
others. Optogenetic studies are a relatively recent advance. 
They are based upon the introduction of light-activated channels 
into specific populations of neurons, permitting relatively precise 
control of action potential generation in live organisms (Zhang et 
al. 2007). For example, Wyart et al. (2009) expressed light 
sensitive genes within so-called Kolmer-Agdur cells of the 
zebrafish, and then non-invasively manipulated the neuronal 
activity of these cells which modulated the swimming behavior 
of the animal.

3. INTERPRETATION OF THE EVIDENCE

We wish to determine if research studies from the categories 
of relational evidence discussed thus far justify the scientific 
claim \( P \), that the brain processes information. Again, we will not 
consider every study; rather, we will start with a typical example 
from each category of evidence and attempt to generalize our 
conclusions to the category.

3.1 S/A

Beginning with Broca, what can we infer from his S/A study of 
subject 'Tan' who could only speak the syllable 'tan', and 
more specifically, how do we clarify the relationship between 
brain structure and the inability to produce fluent, complex 
speech (Broca’s aphasia)? It is clear that Broca’s area lesions 
and Broca’s aphasia are related in the sense that they can occur 
contemporaneously within a single individual. It is also clear 
that Broca’s lesion and Broca’s aphasia need not be 
contemporaneous, for 85% of patients with chronic Broca’s 
aphasia have lesions in Broca’s area, and only 50-60% of 
patients with lesions in Broca’s area have a persisting Broca’s 
aphasia. Further, surgical excision of Broca’s area—a brain 
manipulation/behavioral response study—has led only to 
transient mutism followed by recovery of the patient to normal 
(Dronkers 2000). This evidence alone is enough to at least 
challenge claims of understanding the function of Broca’s area. 
One can of course speculate on the functioning of a brain area 
given an imperfect statistical correlation between that brain area 
and an observed ability, but any claims of knowledge of function 
are excessive: we have knowledge of the correlation and not the 
function.

Let us assume, falsely, that Broca’s lesions and Broca’s 
aphasia were perfectly correlated in the sense that 100% of 
patients with Broca’s aphasia have lesions and 100% of patients 
with lesions have aphasia, for it is possible that other S/A studies 
exhibit perfect correlation. From this finding, can we conclude 
that perfect correlation between structural brain states and 
particular abilities justifies \( P \) ? This would at first appear to 
depend upon the nature of the ability being studied, but it is not 
clear that any S/A study could empirically justify that the brain 
processes information. In S/A studies, one may try to infer brain 
processes given a correlated ability. There is a tendency to argue 
that the structural brain area that is correlated with an ability is in 
fact performing a process proximately responsible for the ability, 
but this is not the logic of S/A studies. The logic of S/A studies 
is as follows: if a subject with structural pattern S cannot do A, 
but a subject without S can do A, then the ability to perform A 
must depend upon S in some way. Even if we accept this logic— 
which requires that the subjects are similar in every other way 
extcept S—we cannot logically infer that the function of S is to 
perform A.

It should be clear that attributing function based upon S/A 
studies is not logically justified. Consider this example. There 
were many times when my computer, the computer I am using 
right now, loses a particular ability that I expect it to have. I 
call a time when I was unable to run programs I typically run, 
and other programs began running very slowly or would shut 
down for no apparent reason in mid-session. The problem turned 
out to be a dead CPU fan. Should we say that the function of the 
CPU fan is to run programs quickly and prevent them from 
shutting down? The abilities in question depended upon the 
spinning of the fan, but the fan does not perform the absent 
abilities—the function of the fan is to cool down the CPU. The 
CPU fan participates in a series of causal interactions that run 
programs when ‘everything is working,’ and we could similarly 
argue, given the results of a S/A study, that a structural feature 
of the brain participates in causal interactions that realize a 
particular ability under certain conditions. This does not entail, 
given a S/A study, that the function of the brain structure is to 
perform the ability in question.

How do we relate the structure of a brain area to the function 
of that area? In S/A studies, that function is always speculatively 
inferred from an observed ability. We do not study how the 
dynamic processes of the brain area and its relations to the rest of 
the organism causally make the ability possible—but this is what 
we need to understand if we are to assign a function (or process) 
to the brain area in question. Broca’s area is more selectively 
related to the ability to produce grammatically appropriate 
speech than some other parts of the organism. One may argue 
that the imperfect but selective correlation between Broca’s area 
and Broca’s aphasia justifies the scientific claim that Broca’s 
aphasia processes linguistic information, but the ability ‘to process 
linguistic information’ plays no obvious role in the study. To 
justify the claim that Broca’s area processes linguistic 
information given the evidence of a related S/A study—a claim 
made but many scientists in the field—we would have to assume 
that speaking grammatically appropriate speech involves 
linguistic information processing by the organism as a whole 
(because this is the only way that processes and operations enter 
into S/A studies. We do not observe brain area processes; we 
observe the functioning organism), and then further identify the 
ability to process linguistic information with the function of 
Broca’s area. But neither of these steps is empirically justified. 
No one directly measures linguistic information processing in the 
study—we simply observe the form of speech—and the 
identification of an organism’s ability with the functioning of a 
brain area is a speculative inference. This does not imply that 
Broca’s area plays no role in the ability to produce fluent
speech—we simply do not know what that role is given limited evidence and theory.

We will not address other S/A studies as the form of our argument applies to all studies that attempt to relate brain structures and organism abilities. In summary, the evidence in S/A studies is composed of correlations between brain structures and organism-level abilities. To arrive at claims of information processing in the brain from S/A studies, one must first infer that the brain structure performs a process or function that directly enacts the organism-level ability in question, but this inference is speculative. In a second layer of interpretation, one must identify performing the process or function with information processing, but this identification enters as an assumption independent of the evidence.

3.2 ES/BR

External-stimulus/brain-response studies address questions of brain function more directly than S/A studies and are probably the largest category of relational evidence. Edgar Adrian, in his pioneering ES/BR research, measured the electrical responses of single sensory cells, such as stretch receptors, while they were fixed to particular weights. He observed that a cell’s electrical responses are in the form of stereotyped action potentials, or spikes, and that the rate of producing spikes increases as the weight increases. Thus the rate, or frequency, of spikes during a fixed time period is able to predict the magnitude of the stimulus.

These early experiments established that single cell responses and stimulus magnitudes may reliably covary with each other. While magnitudes and intensities are important properties of stimuli, they are not the only properties of environmental stimuli that are relevant to an organism. In general, a stimulus may be characterized by multiple properties. For example, an auditory stimulus may be described by its intensity, frequency spectrum, temporal envelope, source direction, source distance, and so on. It is possible that a particular cell responds to one of these properties and not the others, or to some combination of properties, which suggests that a cell may be selective for specific properties or features of the stimulus.

Barlow (1953) was perhaps the first to clearly demonstrate the feature selectivity of sensory cells (Reike et. al 1999). By recording the electrical activity of retinal ganglion cells in the frog, he was able to show that the cell’s activity covaries with the location and size of a circular spot light on the retina. After systematically varying the light spot’s size and location, Barlow determined that the cell’s receptive field—the collection of stimulus properties that maximally activated the cell—is a circularly symmetric form called a center-surround field. Spots of light within a small region of the retina activate the cell, but spots of light away from that region inhibit it.

Hubel and Wiesel (1962) greatly extended Barlow’s work and discovered cells of the striate (visual) cortex that have surprisingly complicated receptive fields. Two of these cell types are the so-called simple and complex cells, which respond maximally to appropriately oriented bars or slits of light. Some of the cells are relatively insensitive to the location of the bar, while others only appreciably respond to moving bars. In describing these cells, Hubel says that “We feel that we have at least some understanding of a cell if we can say that its duty is to take care of a 1 degree by 1 degree region of retina, 6 degrees to the left of the fovea and 4 degrees above it, and to fire whenever a light line on a dark background appears, provided it is inclined at about 45 degrees (Hubel 1962, p168).”

The evidence from all of these pioneering ES/BR electro-physiological studies cannot be interpreted without the concept of selective response. Selective response means, loosely, that the cell fires action potentials only when the ‘right’ stimulus is present. Put more rigorously, selective response refers to two characteristics of neuronal cells: (1) the rate or pattern of firing action potentials (the spike train) covaries with specific stimulus properties, and (2) different cells may respond differently to the same stimulus. Both characteristics are typically implied when referring to the selectivity of cells in ES/BR studies. If someone discovered a neuron that exhibited (1), but on subsequent research discovered that all neurons exhibited (1) in the same way, one would not say that the initial neuron was selective for the stimulus, even though it exhibited selectivity for some stimuli among others. As well, the fact that different neurons respond differently to similar stimuli does not imply (1), since neuronal responses may be random in response to stimuli. Condition (1) is a form of within neuron stimulus selectivity, while condition (2) is a form of between neuron stimulus selectivity. For ES/BR studies such as Hubel and Wiesel’s, when an ES is chosen and controlled by the researcher, we assume that the relation between the ES and BR is causal, as this assumption does not change our interpretation of selectively, even though we use the term ‘covaries’ which has statistical connotations.

We are now in a position to evaluate whether Hubel and Wiesel’s ground-breaking ES/BR studies justify the claim that the brain processes information. In this case we are asking if specific neurons, complex cells of the striate cortex, process or carry information. The experimental evidence consists of recorded responses of complex cells that demonstrate stimulus selectivity in the senses of (1) and (2). It seems that selectivity in the sense of (2) does not provide any justification that complex cells process information; the fact that different cells respond differently to the same stimulus suggests only that the cells are different in some way.

Claims of information processing, if they are justified by this experiment, must follow from the evidence that complex-cell spike trains covary with the properties of visual stimuli, or in causal language, that different visual stimuli cause different complex cell spike trains. Considering the latter causal language, the fact that different causes reliably produce different effects when mediated by the same cell does not appear to justify the claim that the cell processes information, unless one takes that fact to be a definition of information processing itself. Even so, this type of causal relationship appears everywhere one looks. A particular pool ball when hit by other balls with different masses and velocities will undergo different effects. The pool ball may not appreciably move when stimulated by light or sound at typical intensities. The selectivity of the pool ball to acquire
different velocities in response to different causal ‘stimuli’ does not appear fundamentally different than the selectivity of a complex cell, especially if the visual stimulus is taken to be a space-time collection of photons.

On closer analysis, there is a difference between the causality in the pool ball example and the relation between the ES and BR of complex cells. The pool ball example involves direct physical contact and an exchange of energy and momentum, while the causal response of the complex cell is more indirect. Photons travel through the lens of the eye and are absorbed by photoreceptor cells of the retina. Absorption of photons causes the release of the neurotransmitter glutamate at synapses onto so-called bipolar cells, causing the electrical field across the membrane of these cells to become more positive or negative, which respectively increases or decreases the probability of generating an action potential. Bipolar cells have axons that synapse on other cells, and through a series of neuronal connections, influence the membrane potential of complex cells and subsequent action potential generation. The causal chain from photons to complex cell response is complicated and likely includes causal feedback, yet it is not obvious that a complicated causal chain is necessarily information processing.

Even more worrisome is the fact that selective causation need not imply that the BR has any functional relation to the ES at all. Nothing rules out the possibility that those selective correlations are accidental—not in the sense that the correlations are statistically spurious, but that those correlations are functionally irrelevant to the stimuli of interest. As an analogy, suppose my computer has a CPU fan with a blue LED light on the fan. The light, however, is unlit and the fan isn’t spinning. It happens that when I kick my computer just so on the left side of the front cover, the LED lights up, the fan begins spinning but stops after a second or two, and the light goes out. If I kick it again, just so, it starts up for a second then stops. I can reliably cause the fan to turn on for a bit. When I kick the computer in other places, or shake it up, or sing to it, nothing happens to the fan. The fan is selectively correlated with a specific kick. Perhaps there are hundreds of computers, constructed at the same factory, that behave similarly. This selective causal relationship does not imply that the fan is functionally relevant to my kicking, or processes kicking information, or represents kicking. This causal relationship may be accidental. Why then, given the evidence of selective responses in ES/BR studies, do many philosophers and neuroscientists associate information processing with this sort of causation?

3.2.1 Intuition for Information in ES/BR studies

There is a strong tendency to associate information processing with the results of ES/BR experiments like Hubel-Weisel’s. The spike trains of neurons appear to be relaying specific messages about the external environment to the organism. Claude Shannon (1948), the founder of mathematical communication theory, rigorously defined a model of information transfer that may explain this appearance. In Shannon’s language, the physical environment acts as a source that generates a message (ES), the message is transformed by a transmitter—a sensory organ of the organism—into a signal suitable for biological transmission. The spike train (BR) is assumed to be this signal and the neuron to be the transmission channel. These comparisons are reasonable, but the next stage of the communication model, however, is problematic. Communication requires a receiver that performs the inverse operation of the transmitter, or something that reconstructs the environmental message from the spike train signal.

The experimental researcher, the one who discovers selective correlations between neuronal spike trains and environmental messages (stimuli), often plays the surrogate role of the receiver or decoder. By describing relational or mathematical mappings between the ES and BR, neuroscientists attempt to ‘read the neural code.’ But this is not the sort of information transmission we were trying to explain. To complete the biological communication model, and to ground information transfer, we need to explain how the organism can reconstruct the environmental message from its temporal pattern of action potentials, and we must demonstrate that the organism reproduces a similar environmental message within the organism itself. The neuronal spike train is not the message—if anything it is the transmission signal or ‘encoded message.’ Although interesting, it is not enough to show that spike trains have the capacity to represent environmental messages through selective covariation. The fact that researchers can mathematically map spike trains back onto stimuli does not say anything about how the organism physically reconstructs the environmental message. This capacity to map follows immediately from statistical correlations. Neuroscientists who acknowledge these limitations explain that mathematically reconstructing stimuli from spike trains requires taking the homunculus point of view (Reike et al. 1999).

For an organism to receive an environmental message, that message must be within the organism and have the same structure as the original message. This suggestion may appear radical, but it is simply the completion of Shannon’s communication model—the same model that supports the intuition that the brain processes and transmits information. For example, consider telephonic communication. Air pressure waves may be converted into analog electronic messages that are encoded into digital signals and transmitted through a physical channel. This digital signal, which does not mirror the sound wave in form, reaches a destination where it is reconstructed back into an analog message that drives a loudspeaker, reproducing the original pressure wave. If the original message was not reproduced (perhaps imperfectly) at a destination, we could not claim that communication or information transfer took place. A message is communicated if and only if that message is reproduced at the receiver.

If one assumes that the organism receives environmental messages, then in accordance with Shannon’s communication model, at least the structure of that message must be physically reproduced within the organism. The alleged encoded message—or spike train—has a physical basis, thus the message ought to have a physical basis as well. This means that the
scientist would have to demonstrate a set of brain-related physical measurements that copy, perhaps imperfectly, the structure of an environmental message. Let us call this the brain-image of an environmental message. It would remain for the scientist to describe the mechanisms by which neuronal spike trains causally reconstruct the brain-image of a particular environmental message.

But no evidence suggests that brain-images exist, so the very presence of an encoded message within the brain presents a problem. In other words, why should the brain contain encoded messages that transmit environmental messages, yet never reproduce the structure of the message itself? The organism requires the actual message, and not only an encoded version of it. At this point our analogy to Shannon’s communication model breaks down. It does not appear that the environment communicates a message to the organism, but rather, the organism is perhaps translating the environment. Spike trains are not signals corresponding to encoded messages; they are the actual messages only in the language of the organism, whatever that might mean. With respect to the organism, the message is not encoded in anyway, and speaking of a neural code is metaphorical and at times misleading. The analogy has changed from information transfer to language translation. Our goal here, however, is not to support other metaphors, but to show that Shannon’s communication model, which is an integral part of modern technology, does not match the relation between an ES and BR.

Neuroscientists, such as deCharms and Zador (2000), repeatedly claim that spike trains carry information or content about the environment, and suggest what it means ‘to carry information’:

“Imagine recording from the neuron labeled B1 during different types of stimuli or behaviors and discovering the information that this neuron carries about the organism’s environment—the content of this neuron’s signal (p614-15).”

In a concrete example about a retinal cell they say that “The activity of the neuron will be highly correlated with the point of luminance (thus carrying content about this input)(p637).” Like in Hubel-Wiesel’s ES/BR experiments, we call this evidence the selective covariation between stimulus properties and spike trains. deCharms and Zador use the word ‘information’ above to possibly mean ‘specific properties or features of the stimulus.’ Given these examples, we can suppose that they would endorse the following argument: (1) spikes trains and stimulus properties selectively (and causally) covary, and (2) the (representational) content of a spike train is the stimulus property that causes that spike train.

deCharms and Zador do not bring forth any other types of experimental evidence other than selective covariation to justify the claim that spike trains carry informational or representational content, although they do stress that the representational nature of spikes trains is based upon content and function. We have argued that (1) is a statement about the evidence that all of us would agree upon, but that (2) does not obviously follow. The fact that an ES and BR selectively covary, through causal paths, does not appear sufficient to justify claims of representational content, and it has been argued that covariation of this sort is not even necessary for representational content (Millikan 1989, Bechtel 1998).

We need not expect deCharms and Zador, as neuroscientists, to philosophically justify what it means for a spike train to carry informational content, yet if claims of carrying content do not follow immediately from the observed evidence, then we can only assume that they are interpreting the evidence or communicating the evidence by way of metaphor. But deCharms and Zador, along with many other neuroscientists, speak as though ‘carrying content’ is a straightforward experimental fact apart from, or in addition to, selective covariation.

From a philosophical perspective, Dretske (1988, 1995) argues that regular causal covariation, by itself, implies information carrying. For example, he says that flag poles and metal paper clips carry information about temperature because the volumes of these metal objects are reliably correlated with temperature. But is it not too easy to find this sort of information carrying all around us? And why do the objects in question need to be regularly or reliably correlated? Any two things that are causally related, perhaps probabilistically, transmit the same sort of thing. If the flag pole was hit by a lightning bolt, does not the flag pole carry information about the energy of the lightning bolt?

So long as the causal relations are understood between objects c and e, then we might say that e carries information about c. If a situation can be expressed in the form of a law-like equation, then any parameter on one side of the equation can be said to carry information about a parameter on the other side equation, such as the ideal gas law PV=nRT. If the conditions are probabilistic, then we can use probability theory to derive the distribution of one variable given another, so long as we have some understanding of the physical connections between variables. Carrying information, at least according to Dretske, follows directly from knowing the causal relations between two physical situations, or from minimally knowing that two situations are statistically correlated. If by processing information neuroscientists and philosophers mean that stimulus properties causally or statistically covary with regionally specific neuronal activity, then we agree with P, although we suggest abandoning P in favor of more empirically-grounded statements about covariation.

3.2.2 Other Philosophical Justification of Information in ES/BR studies

Considering similar ES/BR experimental evidence, Garson (2003) has attempted to explain a concept of information based upon the pioneering electrophysiological ES/BR studies of Edgar Adrian. Hubel-Wiesel’s and Adrian’s experiments were similar; both consisted of presenting stimuli while measuring the electrical responses of single cells. Although the technologies, organisms, cell types, and stimulus types differed between
implies that a negative change in the frequency of action mapping? And the principle of adaptation is equally troubling; it non-differential mapping as a premise to establish differential specific frequency of action potentials. How can Garson assume Garson’s sense. Rate coding is a form of non-differential principle taken individually contradicts differential mapping in although we are not challenging its methodology. Garson argues—indeed, his belief in differential mapping, which he takes from Adrian’s work, does not correspond to the predominant experimental methodology used to acquire evidence in ES/BR studies. Since Hubel-Wiesel’s experiments, neuronal responses in electrophysiological ES/BR studies are most often understood by characterizing the feature selectivity of the cell type (Rieke et al., p12). This selectively corresponds to the collection of stimulus properties that evoke responses for that cell, and highlights the properties that evoke optimal responses. It is based upon the concept of selective response that we analyzed above. Selectivity involves a non-differential mapping between stimuli and responses, it grounds our current understanding of sensory cell types, and directly opposes Garson’s concept of differential mapping.

But the fact that a scientific community makes use of feature selectivity rather than differential mapping to acquire evidence does not in itself deny differential mapping. We, too, are questioning the community in its interpretations of evidence, although we are not challenging its methodology. Garson argues that the differential mapping between stimuli and spike rates logically follows from a conjunction of the principle of neuronal rate coding with the principle of adaptation:

“While the principle of rate coding entails that the frequency of the sequence of action potentials is an exponential function of the magnitude of the stimulus, the principle of adaptation entails that upon application of a constant stimulus, the frequency of the sequence of action potentials will diminish, and eventually such outputs will stop being produced. Hence the relation between the sequence and stimulus is differential.”

No further derivation is given, which is concerning since each principle taken individually contradicts differential mapping in Garson’s sense. Rate coding is a form of non-differential mapping: a specific stimulus intensity directly maps onto a specific frequency of action potentials. How can Garson assume non-differential mapping as a premise to establish differential mapping? And the principle of adaptation is equally troubling; it implies that a negative change in the frequency of action potentials maps onto a constant intensity of the stimulus. But this directly conflicts with Garson’s claim that differences in spike rates map onto differences in intensity, and “not to a constant state of the stimulus.”

We agree with Garson that differences in stimuli are particularly important to the human organism and other animals, but Garson does not logically establish differential mapping—or does the scientific evidence primarily support differential mapping—and thus he does reach the goal of deriving a concept of information from the evidence. Even if he did establish differential mapping, the next step in the derivation, where he argues that the relation between stimuli and spike trains is arbitrary, meets with difficulties as we will show.

Garson provides an alternative way to understand the relation between stimuli and spike rates that does not directly depend upon the truth of differential mapping. He explains that a given stimulus s1 may be associated with multiple firing rates r1 and r2 because of adaptation, and that two stimuli s1 and s2 may be associated with a single firing rate r1 for similar reasons. In his words:

“For example, suppose a stimulus of intensity s1 elicits a firing rate ri from a neuron. Then s1 is held constant, and by the principle of adaptation, the firing rate is reduced (say, to r1,2). Upon increasing the stimulus to s1+1, the firing rate may return to r1, its initial value.”

Garson is arguing that the relation between stimuli and firing rates is a many-to-many relation as opposed to a one-to-one relation or a one-to-many relation. This sense of relation corresponds to the use of relation in relational databases. For example, the relation between a person and that person’s driver’s license number is one-to-one; the relation between each continent and its countries is one-to-many—each continent can have multiple countries, but each country belongs to one continent; and the relation between academic articles and their authors is many-to-many—each article may have multiple authors, and each author may contribute to multiple articles.

Garson believes that the many-to-many relation between stimuli and spike rates supports a notion of arbitrariness, and that this arbitrariness, although not sufficient, is a necessary constituent of a concept of information. But what is necessarily arbitrary about a many-to-many relation in contrast to a one-to-one relation? A one-to-one relation, such as the driver’s license example above, may be as arbitrary—or more arbitrary—than a many-to-many relation, depending upon how that relation came to be. Garson attempts to distinguish between arbitrary and non-arbitrary causal relations by contrasting two neurophysiological mechanisms involved in auditory perception: the tympanic membrane and the hair cells of the inner ear. He says that the relation between air pressure oscillations and the vibrations of the tympanic member is one-to-one, and is therefore non-arbitrary; but that the hair cells of the inner ear undergo adaptation, which implies a many-to-many relation between air pressure oscillations and hair cell firing, and is therefore arbitrary. Garson believes that relations that come to be through mechanisms of adaptation and are many-to-many are arbitrary, and those that do not involve adaptation and are one-to-one are
non-arbitrary. But it seems obvious that one-to-one and many-to-many relations may both be arbitrary or non-arbitrary. The relevant difference that determines arbitrariness for Garson must be the presence of adaptive mechanisms, but he uses neuronal adaptation only as a means to justify the many-to-many relation between stimuli and responses. In this sense the argument is circular.

3.2.3 Summary of ES/BR Studies

Rigorous experimental neuroscience has demonstrated neurons that selectively respond to a wide range of measurable parameters across the five senses. Because selective responses reliably covary with stimulus properties, we can use mathematical tools to predict stimulus properties given spike trains, and predict spike trains given stimulus properties. The idea that there is a neural code, however, is a metaphor, since spike trains are not encoded with respect to the organism, anymore than English is encoded with respect to an English speaker. Further, the idea that spikes trains carry information is grounded in the experimental evidence of selective covariations, but most commentators conflate or equate carrying information and selective covariation.

3.3 T/BR

Task/brain-response studies combine aspects of S/A and ES/BR studies, and our critiques of these studies will apply. In addition to systematically manipulating the external environment as in ES/BR studies, T/BR studies add to this manipulation a task for the subject to perform. The task is similar to an ability in a S/A study, except that the task is a transient activity while an ability is an ongoing capacity to act in a particular way. With regard to scientific research, tasks should have observable or measurable criteria for successful completion, while abilities should have observable or measurable criteria for possession of the ability. We can often study a topic using either task or ability language. For instance, in memory studies, we can assign subjects the task of memorizing a set or numbers, and then ask for those numbers at a later time. The task of remembering and the ability to remember are similar in that the criteria for completion of the task and possession of the ability are equivalent. If one remembers the numbers correctly, one has successfully completed the task and possesses the ability to remember.

Let us recall the Poppenk et al. (2010) T/BR study on prospective memory described above, where subjects were presented a series of visual scenes and instructed to either imagine performing an action associated with that visual scene (e.g. swinging on a swing when shown a swing), or to use the scene as a reminder to perform an action the next time the same scene was viewed. FMRI was used to measure brain properties while the subjects performed these tasks. After this task, the subjects were taken to a quite room to perform an identification test. They were shown visual scenes on a computer and asked to indicate whether each scene was studied as an intention, an action, or not seen during scanning at all. Researchers recorded correct and incorrect responses. The results of the identification test were statistically correlated with the fMRI data to identify spatiotemporal fMRI activity patterns that predicted correct responses on the identification test. Poppenk et al. speculated that some of the identified brain regions enact “processes associated with successful encoding of intentions (p911).”

This particular T/BR study is more complicated than the ES/BR studies of Hubel-Wiesel from an interpretational standpoint. Although T/BR studies are not necessarily more complicated than ES/BR studies, the complexity of Poppenk et al.’s study is not atypical for fMRI studies that include cognitive tasks. Like S/A and ES/BR studies described above, the empirical evidence in this T/BR study consists of selective correlations, in this case between successful task completion and properties of brain areas. These selective correlations, like those described in S/A and ES/BR studies, do not logically imply that the function of the identified brain regions is to perform a process directly related to the task. Poppenk et al. make no attempt to understand the so-called processes of the identified brain region other than to say that processes are associated with the task, but the observed form of this association is statistical correlation. Even if this association was selectively causal, we still could not infer that the function of the brain region had to do with completing the task, for the causal association could be accidental. And even if the function of the brain region involves processes to complete the task, we do not know that completing the task involves processing information of any kind.

3.4 BM/BR

Brain-manipulation/behavioral response studies demonstrate the behavioral effects of causally manipulating brain properties. As an example, optogenetic studies are a relatively recent advance in BM/BR experimentation, and permit precise manipulation of neuronal activity. These experiments involve expressing light-sensitive genes within specific neurons or populations of neurons in living animals. When the neurons with the expressed genes are exposed to light of a particular wavelength, the activity of the neuron will either increase or decrease, allowing for precise control of the neuron’s activity. Presumably the expressed genes do not significantly alter the functioning of the neuron otherwise. For example, Wyart et al. (2009) expressed light sensitive genes within so-called Kolmer-Agdur cells of the zebrafish, and then non-invasively manipulated the neuronal activity of these cells which modulated the swimming behavior of the animal.

BM/BR studies establish causal relationships between the activity of multiple brain areas or between brain area activity and behavioral responses. Canonical examples of BM/BR experiments involve electrical stimulation of brain areas resulting in muscle movements. These types of experiments can be traced back to at least Fritsch and Hitzig (1870) who applied surface electrodes to dog brain and demonstrated that the anatomical location of electrical stimulation selectively covaried with movements in different muscle groups. Neuroscientists name the
structure of this covariation a somatotopic organization, and classically explain the relation between cerebral cortex and muscle movements with the following three hypotheses (Graziano et al. 2002): (1) the precentral gyrus, or primary motor cortex, contains an explicit topographic map of the body with the foot on the top of the cerebral hemisphere, the mouth on the bottom, and other parts systematically organized in between; (2) the activity at each point in the map specifies the tension in a small group of muscle fibers; and (3) cortical motor areas are organized in a clear hierarchy of control. These classical hypotheses have been summarized as far back as 1938 by Fulton, although there is a significant body of subsequent evidence that is not compatible with the classical theory (Graziano et al. 2002).

Contemporary BM/BR studies of primate motor function typically involve inserting microelectrodes into the cerebral cortex of an awake animal, and injecting low electrical currents into the cellular network while simultaneously recording the pattern of muscle movements, allowing the researcher to catalog the causal relations between electrical stimulation and these movements. One hopes or assumes that the patterns of muscle movements in response to exogenous stimulation are similar to those caused by endogenous neural activity, although the induced electrical activity is clearly non-physiologic, complicating any interpretation of the results.

If we consider the brain to be a mechanical mechanism, what do these motor BM/BR studies tell us about the brain? In other words, knowing that anatomical locations of electrical stimulation and patterns of muscle movement covary with each other, what can we say about the processes or functions that occur in those brain areas? We might say that there is a causal propagation of electrical activity, beginning from motor cortical areas through the central nervous system and to spinal motor nerves that enact patterns of muscle contraction. If we electrically stimulate the brain in other areas, the electrical activity does not propagate to spinal motor nerves. The motor cortex therefore acts as a metaphorical gateway or hub of electrical activity from CNS to spinal nerves, where the pathways are at least partially organized with respect to specific muscle movements.

Claims of motor information processing are not needed to describe these results, but presumably arise when one assumes that areas of the motor cortex naturally represent various muscle groups, or that the precentral gyrus contains a map of the body, but the fact that electrical activity propagates through specific pathways does not establish that motor areas naturally represent muscle groups. Just as the idea of a neural code is a sometimes useful metaphor for communicating scientific results, the idea of a topographic map of the body in the precentral gyrus is a useful metaphor for summarizing the data about causal organization with respect to electrical stimulation.

Compare the structure of the patellar reflex in the peripheral nervous system: hitting the patellar tendon stretches the quadriceps muscle which activates sensory receptors that propagate electrical activity through motor neurons that contract the quadriceps. We can describe the propagation of physical changes without any reference to information processing or transmission. Of course, one can use a metaphorical information language to describe the propagation of electrical activity involving the patellar reflex, but this language adds nothing to our physical understanding of the reflex. Electrical activity does not propagate randomly through the brain; its pathways are organized, and scientists attempt to understand this causal organization with respect to an organism’s abilities. BM/BR studies are an important tool for understanding this organization, however, knowledge of causal organization does not imply knowledge of the processes that occur within a brain region, nor does it provide evidence of information processing.

### 4. CONCLUSIONS

Cognitive and neuroscientists have measured action potentials, ischemic lesions, distributions of dopamine receptors, electromagnetic surface potentials, and many other brain properties. They have measured temporal sequences of brain properties contemporaneous with environmental stimuli and behavioral actions, and have manipulated brain tissue and catalogued behavioral changes. It is clear that the activity of particular brain areas selectively covaries with specific stimulus properties and abilities. However, since we cannot logically infer processes and functions given experimental correlations between stimuli or abilities and brain areas, it is difficult to justify Bechtel and Richardson’s supposed uncontroversial claim that “the brain contains some regions that are specialized for processing specific types of information.” The brain does contain regions that selectively covary with particular stimulus properties and particular abilities, and we may at times attribute causality to these covariations, but interpretational claims that attribute information processing to brain areas go beyond the evidence.

We admit that our examination of the research is far from exhaustive, and that we have limited our analysis to relational evidence. There are likely other categories of evidence that support information processing in the brain, but we feel we have examined evidence that is classically taken to support information processing and find it inadequate.

Rather than inferring solely from the experimental evidence as we have suggested, cognitive scientists typically begin with a priori theories about particular cognitive processes, and use the results of relational studies to choose between these theoretical cognitive processes (Henson 2006). One might call the claim that the brain processes information one of these hypothetical cognitive theories. We have not found empirical support for this cognitive theory, although others may interpret the evidence otherwise. If one assumes that selective correlation or causation implies information carrying, then one will see information processing everywhere one looks. In this sense, information processing is not a scientific theory or fact, but a basic principle or metaphor that many people find useful in communicating and interpreting evidence.

There is no doubt that mathematical modeling will help us to understand the functioning of the brain, but we have not
addressed modeling efforts in this work because modeling in
cognitive science is underdetermined, and to our knowledge
there are no cognitive models that are largely accepted as
representing mental facts. Further, while modeling may provide
some evidence for information processing in the brain, this
evidence is relatively weak because of the above and because
information processing enters as an assumption of these models
and not as a consequence.

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