Functional brain imaging with use of a new and powerful neuroimaging technique

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ABSTRACT

Most of the information available on the human brain came from subjects who had sustained major head wounds, or who suffered from various mental disorders. By determining the extent of brain damage, and the nature of the loss of function, it was possible to infer which regions of the brain were responsible for which function. With the development of the imaging techniques of computerised tomography (CT) and magnetic resonance imaging it was possible to be more specific as to the location of damage in brain injured patients. The measurement of the electrical signals on the scalp, arising from the synchronous firing of the neurons in response to a stimulus, known as electroencephalography (EEG), opened up new possibilities in studying brain function in normal subjects. However it was the advent of the functional imaging modalities of positron emission tomography (PET), single photon emission computed tomography (SPECT), functional magnetic resonance imaging (fMRI), and magnetoencephalography (MEG) that led to a new era in the study of brain function. In this paper the mechanisms of the techniques mentioned above are outlined, together with an assessment of their strengths and weaknesses. Then an introduction to the Metabolism and Blood Flow in the Brain is given. This is followed by a more detailed explanation of functional MRI and how such experiments are performed.

Keywords: Functional Magnetic Resonance Imaging; Brain Function; EEG

1. INTRODUCTION

Functional brain imaging using MRI (functional MRI or fMRI) has become a valuable tool for studying function/structure relationships in the human brain in both normal and clinical populations. This paper describes the physiological changes associated brain with activity, including changes in blood flow, volume, and oxygenation. The latter of these, known as Blood Oxygenation Level Depended (BOLD) contrast, is the most common approach for functional MRI, but it is related to brain activity via a variety of complex mechanisms. Blood oxyg enation level dependent functional MRI and near infrared optical tomodraphy have been widely used to investigate hemodynamic responses to functional stimulation in the human brain. The temporal hemodynamic response shows an increased total hemoglobin concentration, which indicates an increased cerebral blood volume (CBV) during physiological activation. Blood Oxygenation Level Dependent signal indirect measure of neural activity. The signal variations induced by respiration and cardiac motion decrease the statistical significance in functional MRI data analysis. Physiological noise can be estimated and removed adaptively using signal projecting technique with the actual functional signal preserved. We estimate and remove the physiological noise from the magnitude images. This method is a fully data-driven method, which can efficiently and effectively reduce the overall signal fluctuation of functional MRI data. Assumes that the MRI data recorded on each trial are composed of a signal added with noise. Signal (random) is present on every trial, so it remains constant through averaging and Noise randomly varies across trials, so it decreases with averaging. Thus, Signal-to-Noise Ratio (SNR) increases with averaging.

2. COMPARISON OF THE FUNCTIONAL BRAIN IMAGING MODALITIES

The brain imaging techniques that had been presented in this paper all measure slightly different properties of the brain as it carries out cognitive tasks. Because of this the techniques should be seen as complementary rather than competitive. All of them have the potential to reveal much about the function of the brain and they will no doubt develop in clinical usefulness as more about the underlying mechanisms of each are understood, and the hardware becomes more available. A summary of the
strengths and weaknesses of the techniques is presented in Table 1 [5].

2.1. SPECT and PET

The imaging modalities of single photon emission computed tomography (SPECT) and positron emission tomography (PET) both involve the use of radioactive nuclides either from natural or synthetic sources. Their strength is in the fact that, since the radioactivity is introduced, they can be used in tracer studies where a radiopharmaceutical is selectively absorbed in a region of the brain. The main aim of SPECT as used in brain imaging, is to measure the regional cerebral blood flow (rCBF). The earliest experiments to measure cerebral blood flow were performed in 1948 by Kety and Schmidt [1]. They used nitrous oxide as an indicator in the blood, measuring the differences between the arterial input and venous outflow, from which the cellular uptake could be determined.

This could only be used to measure the global cerebral blood flow, and so in 1963 Glass and Harper[2], building on the work of Ingvar and Lassen, used the radioisotope Xe-133, which emits gamma rays, to measure the regional cerebral blood flow. The development of computed tomodraphy in the 1970’s allowed mapping of the distribution of the radioisotopes in the brain, and led to the technique now called SPECT [3].

2.2. EEG and MEG

Measuring the electrical signals from the brain has been carried out for several decades [4], but it is only more recently that attempts have been made to map electrical and magnetic activity. The electroencephalogram is recorded using electrodes, usually silver coated with silver chloride, attached to the scalp and kept in good electrical contact using conductive electrode jelly. One or more active sites may be monitored relative to a reference electrode placed on an area of low response activity such as the earlobe. The signals are of the order of 50 microvolts, and so care must be taken to reduce interference from external sources, eye movement and muscle activity. Several characteristic frequencies are detected in the human EEG. For example, when the subject is relaxed the EEG consists mainly of frequencies in the range 8 to 13 Hz, called alpha waves, but when the subject is more alert the frequencies detected in the signal rise above 13Hz, called beta waves. Measurements of the EEG during sleep have revealed periods of high frequency waves, known as rapid eye movement (REM) sleep which has been associated with dreaming [5].

MEG experiments are carried out in much the same way as their EEG counterpart. Having identified the peak of interest, the signals from all the detectors are analysed to obtain a field map. From this map an attempt can be made to ascertain the source of the signal by solving the inverse problem. Since the inverse problem has no unique solution, assumptions need to be made, but providing there are only a few activated sites, close to the scalp then relatively accurate localization is possible, giving a resolution of the order of a few millimeters.

MEG has the advantage over EEG that signal localisation is, to an extent, possible, and over PET and fMRI in that it has excellent temporal resolution of neuronal events. However MEG is costly and its ability to accurately detect events in deeper brain structures is limited.

2.3. Functional MRI and MRS

Since functional magnetic resonance imaging is the subject of this paper, little will be said in this section as to the mechanisms and applications of the technique. The purpose of this section is to compare fMRI to the other modalities already mentioned, and also to consider the related, but distinct technique of magnetic resonance spectroscopy (MRS) [5].

During an fMRI experiment, the brain of the subject is scanned repeatedly, usually using the fast imaging technique of echo planar imaging (EPI). The subject is required to carry out some task consisting of periods of activity and periods of rest. During the activity, the MR signal from the region of the brain involved in the task normally increases due to the flow of oxygenated blood into that region. Signal processing is then used to reveal these regions. The main advantage of MRI over its closest counterpart, PET, is that it requires no contrast agent to be administered, and so is considerably safer. In addition, high quality anatomical images can be obtained in the same session as the functional studies, giving greater confidence as to the source of the activation. However, the function that is mapped is based on blood flow, and it is not yet possible to directly map neuroreceptors as PET can. The technique is relatively expensive, although comparable with PET, however since many hospitals now have an MRI scanner the availability of the technique is more widespread [5].

Table 1. Comparison of modalities for studying brain function.

<table>
<thead>
<tr>
<th>Technique</th>
<th>Resolution</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>SPECT</td>
<td>10 mm</td>
<td>Low cost</td>
<td>Invasive, Limited resolution</td>
</tr>
<tr>
<td>PET</td>
<td>5 mm</td>
<td>Sensitive, Good resolution, Metabolic studies, Receptor mapping</td>
<td>Invasive, Very expensive</td>
</tr>
<tr>
<td>EEG</td>
<td>poor</td>
<td>Very low cost, Sleep and operation monitoring</td>
<td>Not an imaging technique</td>
</tr>
<tr>
<td>MEG</td>
<td>5 mm</td>
<td>High temporal resolution</td>
<td>Expensive, Limited resolution for deep structures</td>
</tr>
<tr>
<td>fMRI</td>
<td>3 mm</td>
<td>Excellent resolution, Non-invasive</td>
<td>Expensive, Limited to activation studies</td>
</tr>
<tr>
<td>MRS</td>
<td>low</td>
<td>Non-invasive metabolic studies</td>
<td>Expensive, Low resolution</td>
</tr>
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FMRI is limited to activation studies, which it performs with good spatial resolution. If the resolution is reduced somewhat then it is also possible to carry out spectroscopy, which is chemically specific, and can follow many metabolic processes. Since fMRS can give the rate of glucose utilisation, it provides useful additional information to the blood flow and oxygenation measurements from FMRI, in the study of brain metabolism [5].

3. METABOLISM AND BLOOD FLOW IN THE BRAIN

The biochemical reactions that transmit neural information via action potentials and neurotransmitters, all require energy.

This energy is provided in the form of ATP, which is produced from glucose by oxidative phosphorylation and the Kreb's cycle (Figure 1) [5].

As ATP is hydrolysed to ADP, energy is given up, which can be used to drive biochemical reactions that require free energy. The production of ATP from ADP by oxidative phosphorylation is governed by demand, so that the energy reserves are kept constant. That is to say, the rate of this reaction depends mainly on the level of ADP present.

This means that the rate of oxygen consumption by oxidative phosphorylation is a good measure of the rate of use of energy in that area [5].

The oxygen required by metabolism is supplied in the blood. Since oxygen is not very soluble in water, the blood contains a protein that oxygen can bind to, called haemoglobin. The important part of the haemoglobin molecule is an iron atom, bound in an organic structure, and it is this iron atom which gives blood its colour. When an oxygen molecule binds to haemoglobin, it is said to be oxyhaemoglobin, and when no oxygen is bound it is called deoxyhaemoglobin.

To keep up with the high energy demand of the brain, oxygen delivery and blood flow to this organ is quite large. Although the brain's weight is only 2% of the body’s, its oxygen consumption rate is 20% of the body’s and blood flow 15%. The blood flow to the grey matter, which is a synapse rich area, is about 10 times that to the white matter per unit volume. Regulation of the regional blood flow is poorly understood, but it is known that localised neural activity results in a rapid selective increase in blood flow to that area [5].

4. BLOOD OXYGEN LEVEL DEPENDENT CONTRAST IN MR IMAGES

Since regional blood flow is closely related to neural activity, measurement of the rCBF is useful in studying brain function. It is possible to measure blood perfusion with MRI, using techniques similar to those mentioned. However there is another, more sensitive, contrast mechanism which depends on the blood oxygenation level, known as blood oxygen level dependent (BOLD) contrast. The mechanisms behind the BOLD contrast are still to be determined, however there are hypotheses to explain the observed signal changes.

Deoxyhaemoglobin is a paramagnetic molecule whereas oxyhaemoglobin is diamagnetic. The presence of deoxyhaemoglobin in a blood vessel causes a susceptibility difference between the vessel and its surrounding tissue. Such susceptibility differences cause dephasing of the MR proton signal [6], leading to a reduction in the value of T2*. In a T2* weighted imaging experiment, the presence of deoxyhaemoglobin in the blood vessels [7, 8] causes a darkening of the image in those voxels containing vessels. Since oxyhaemoglobin is diamagnetic and does not produce the same dephasing, changes in oxygenation of the blood can be observed as the signal changes in T2* weighted images [5].

It would be expected that upon neural activity, since oxygen consumption is increased, that the level of deoxyhaemoglobin in the blood would also increase, and the MR signal would decrease. However what is observed is an increase in signal, implying a decrease in deoxyhaemoglobin. This is because upon neural activity, as well as the slight increase in oxygen extraction from the blood, there is a much larger increase in cerebral blood flow, bringing with it more oxyhaemoglobin (Figure 2). Thus the bulk effect upon neural activity is a regional decrease in paramagnetic deoxyhaemoglobin, and an increase in signal [5].

The study of these mechanisms are helped by results from PET and near-infrared spectroscopy (NIRS) studies. PET has shown that changes in cerebral blood flow and cerebral blood volume upon activation, are not accompanied by any significant increase in tissue oxygen consumption [9].

NIRS can measure the changes in concentrations of oxy- and deoxyhaemoglobin, by looking at the absorbency at different frequencies. Such studies have shown an increase in oxyhaemoglobin, and a decrease deoxyhaemoglobin upon activation. An increase in the total amount of haemoglobin is also observed, reflecting the increase in blood volume upon activation [2].

Figure 1. Overview of the aerobic metabolism of glucose to ATP following the Kreb’s cycle.
the major optimization goals in fMRI and several fMRI acquisition approaches. Substantial progress is being made to reduce artifacts in fMRI as well as to improve the measurement of alternate physiological phenomena using MRI.

The spatial activation pattern of changes in deoxyhemoglobin concentration is consistent with the BOLD signal map. The patterns of oxy- and deoxyhemoglobin concentrations are very similar to one another. The temporal hemodynamic response shows an increased total hemoglobin concentration, which indicates an increment of CBV during physiological activation. It has now been firmly established that magnetic resonance imaging can be used to map brain function.

The main impetus of research and development of the technique, needs to be directed in several areas if fMRI is to become more than 'colour phrenology', intriguing in its results yet having little clinical value. The mechanisms behind the BOLD effect need to be better understood, as does the physiological basis of the observed blood flow and oxygenation changes. The combination of the functional imaging modalities needs attention, since it is unlikely that any one method will provide the full picture. Finally, robust and simple techniques for data analysis need to be developed, allowing those who do not specialise in fMRI, to carry out experiments and interpret results.

5. CONCLUSION

Functional magnetic resonance imaging can accurately represent cerebral topography, cortical venous structures and underlying lesions. Functional activation appears to accurately localize appropriate cortical areas and these studies are feasible in the presence of local pathology. It is extremely useful in presurgical planning as well as assessment of operability. Intra-operatively, it shows a great promise in being able to define the exact location and extent of lesions with respect to surrounding functional cortex.

Functional MRI is a new and powerful neuroimaging technique that can create an anatomical and functional model of an individual patient's brain. The concurrent 3-D rendering of cerebral topography, cortical veins and related pathology gives an unprecedented display of critical relational anatomy. Since stereotaxy means simply the three dimensional arrangement of objects, then fMRI may be the ultimate stereotactic system. It allows us to see through the scalp and cortex into subcortical areas which are not visually apparent. It accurately predicts cortical gyral and venous anatomy as well as the subcortical location and extent of lesions. But most importantly, it is capable of mapping specific cortical functions to anatomical regions thereby combining form and function.

We have described the physiological bases of functional MRI and introduced a physiologically relevant model of the vascular response to fMRI. We described

REFERENCES