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Noninvasive techniques for probing neurocircuitry and treating illness: vagus nerve stimulation (VNS), transcranial magnetic stimulation (TMS) and transcranial direct current stimulation (tDCS)

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Abstract

Although the preceding chapters discuss much of the new knowledge of neurocircuitry of neuropsychiatric diseases, and an invasive approach to treatment, this chapter describes and reviews the noninvasive methods of testing circuit-based theories and treating neuropsychiatric diseases that do not involve implanting electrodes into the brain or on its surface. These techniques are transcranial magnetic stimulation, vagus nerve stimulation, and transcranial direct current stimulation. Two of these approaches have FDA approval as therapies.

Keywords:

TMS, VNS, neuromodulation

INTRODUCTION

The techniques discussed in this chapter serve as an appropriate last chapter for this volume on brain circuits. The preceding chapters have outlined the emerging theories about circuit abnormalities in neuropsychiatric disorders. The chapter on deep brain stimulation (DBS) reviews the most invasive method for stimulating and modifying the behavior within these circuits. In contrast, the technologies in this chapter are 'noninvasive'. We recognize that some would argue that electrically stimulating the brain is always invasive, and that many, maybe even all, brain stimulation techniques are not noninvasive. This is especially true of techniques that require general anesthesia and the implantation of a permanent pacemaker such as vagus nerve stimulation (VNS). However, for the pedagogical and nosological purposes of separating this chapter from the preceding, the word 'noninvasive' means a brain stimulation method that does not involve craniotomy and implantation of an electrode into the brain (eg DBS or resting on the dura (epidural cortical stimulation)). Thus, the current 'noninvasive' techniques with either existing or promising clinical applications are, in declining order of invasiveness, VNS, transcranial magnetic stimulation (TMS), and transcranial direct current stimulation (tDCS). For each of these technologies, we briefly describe the technique, discuss the major ideas concerning mechanisms of action, and then touch on safety. Following this we overview the research and clinical uses. There are now entire journals devoted to the field of brain stimulation (Sackeim and George, 2008), and entire books devoted to each of the individual techniques (George and Belmaker, 2006) as well as in-depth overviews (Higgins and

<u>George</u>, 2008). The interested reader is referred to these. This chapter will focus on a quick and precise introduction to each with a focus on emerging clinical applications and research uses pertinent to the preceding chapters and theories.

One of the recurring themes within each of the techniques is the currently inadequate understanding of the translational neurobiological effects of the 'use parameters'. These are the pulse width, current direction, intensity, frequency, duty cycle, and the overall dose as well as dosing scheme. The future of the promising field of brain stimulation will undoubtedly involve better translating the knowledge gained about appropriate use parameters from preclinical cellular and nonhuman animal studies into clinical brain stimulation therapeutic uses.

VAGUS NERVE STIMULATION

The idea of stimulating the vagus nerve to modify central brain activity has been pursued for over 100 years. However, it was not until the mid-1980s that methods became available to efficiently stimulate the vagus in man and animals.

Description of Method

Although one can stimulate the vagus in several different ways, even transcutaneously, for all intents and purposes VNS in the modern literature refers to a technique where a surgeon (for human studies) or researcher (for nonhuman animal uses) wraps a unidirectional wire around the vagus nerve in the neck (see Figure 1). This wire is then connected to a subcutaneous, battery operated, generator, which is implanted subcutaneously in the left chest wall, which intermittently sends an electrical current through the wire and thus through the nerve that then conveys a signal through neural impulses into the brainstem (George *et al*, 2000).

Figure 1.



Clinical vagus nerve stimulation (VNS). The VNS generator (a) contains a small battery that generates electrical impulses. A surgeon implants the generator subcutaneously over the chest (b) and attaches the electrodes to the left vagus nerve (c). Intermittent signals from the VNS device travel up the vagus nerve (d) and enter the medulla. (Reprinted with permission from APPI, from <u>Higgins and George (2008</u>)).

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VNS implantation is usually an outpatient procedure in the United States typically preformed (sic) by neurosurgeons. The battery in the device generates an intermittent electrical stimulation that is delivered to the vagus nerve. Clinicians following the patient control the frequency and intensity of the stimulation. Adjustments to the stimulation parameters are transmitted from a computer to the VNS device by a handheld infrared wand placed over the device.

The stimulating wire wrapped around the nerve is directional, and this unidirectional feature likely helps minimize efferent side effects from stimulating vagal efferent (descending) fibers. However, it is likely that at least some patients have had the leads reversed, without noticeable harm (Koo *et al*, 2001).

The vagus nerve is actually a large nerve bundle, composed of different sized nerves (both unmyelinated and myelinated). The vagus nerve is thus a complex structure and the current form of VNS is imprecise with respect to activating discrete nerves within the bundle. Microsurgical techniques might theoretically allow for more focal VNS.

Putative Mechanisms of Action

To refresh, the vagus nerve (10th cranial nerve) enters the brain at the medulla. It is the longest cranial nerve extending into the chest and abdominal cavity. 'Vagus' comes from the Latin word for wandering, and this nerve is remarkably complex, both in where it comes from, and the variety of information it passes bidirectionally between the brain and the viscera. Traditionally, the vagus nerve has been conceptualized as modulating the parasympathetic tone of the internal organs (efferent functions). However, 80% of the signals traveling through the vagus nerve actually go from the organs back into the brain (afferent) (Foley and DuBois, 1937).

In 1938, Bailey and Bremer (Bailey and Bremer, 1938) stimulated the vagus nerve of cats and reported that this synchronized the electrical activity in the orbital cortex. In 1949, Paul MacLean and Karl Pribram carried out similar studies with anesthetized monkeys. Using electroencephalogram (EEG) recordings they found that VNS generated slow waves over the lateral frontal cortex (Maclean, 1990). The afferent fibers traveling in the vagus terminate largely in the nucleus tractus solitarius (NTS) in the medulla. The NTS, in turn, innervates the noradrenergic nucleus locus coeruleus (LC) directly (Van Bockstaele et al, 1999a, 1999b) as well as indirectly through the rostral ventrolateral medulla (Van Bockstaele *et al*, 1989), which sends strong projections to LC neurons (Aston-Jones *et al*, 1986; Ennis and Aston-Jones, 1988). LC neurons project extensively throughout the neuraxis, providing prominent noradrenergic innervation in the orbitofrontal cortex and the insula, including somatotopically defined regions that may represent emotional (limbic) information (Aston-Jones, 2004). Thus, it is plausible that the NTS regulates NE release in the forebrain through its descending projections to LC afferents in the PGi. In addition, these connections show that many vagus afferent fibers connect transynaptically to areas of the limbic brain that regulate emotion. It is no surprise then that when we grieve we have the perception of having a 'broken heart', or feel like there are 'butterflies in our stomach' when we are nervous or anxious. This misplacement concerning the source of the sensory signal may reflect the fact that vagal cardiac fibers terminate in brain regions where the limbic system and gut sensations overlap.

Jake Zabara in the mid-1980s was perhaps the first to show convincingly the therapeutic benefits of VNS, although many had been considering this avenue before Zabara (<u>Groves and Brown, 2005</u>). Zabara discovered in a canine model of epilepsy (strychnine induced) that repetitive electrical stimulation of the vagus nerve was able to acutely terminate a motor seizure. Importantly, he also found that the anticonvulsant benefits could outlast the period of stimulation by a factor of four (<u>Zabara, 1985b</u>, <u>1992</u>). Constant stimulation was not required for enduring anticonvulsive effects.

Safety (regarding a device implant- not transcutaneous stimulation)

The adverse events associated with VNS fall into two categories—those associated with the complications of the surgery and those resulting from the side effects of stimulation. The risks associated with surgery are minimal ($O'Reardon \ et \ al, \ 2006$). Wound infections are infrequent (less than 3%) and managed with antibiotics. Pain at the surgical site almost always resolves within 2 weeks. Rarely left vocal cord paresis persists after surgery (<1 in 1000), but usually resolves slowly over the ensuing weeks.

Temporary asystole during the initial testing of the device is a rare but serious surgical complication. In approximately 1 out of 1000 cases asystole has been reported in the operating room during initial lead testing. It may be a result of aberrant electrical stimulation resulting from poor hemostatic control. That is, blood in the surgical field causes arcing of the current and the cardiac branch gets depolarized. Fortunately, no deaths have been reported as normal cardiac rhythm has always been restored. Postoperatively these patients have been able to safely use VNS. More importantly and surprisingly given the known efferent VNS effects, no cardiac events have been reported when the device is turned on for the first time after surgery.

The most common side effects associated with stimulation are hoarseness, dyspnea, and cough. They are dose dependent and correlate with stimulation intensity and can be minimized with reductions in the stimulation parameters. Interestingly, most side effects decrease with time (<u>Sackeim *et al*</u>, 2001c). Hoarseness or voice alteration is the most persistent problem. Between 30 and 60% continue to experience this side effect during times of stimulation although for reasons that are unclear this also

diminishes over months to years. One would speculate that VNS might induce a parasympathetic response. However, this has been aggressively monitored and has not been an issue.

VNS therapy also affects respiration during sleep and has been shown to worsen preexisting obstructive sleep apnea/hypopnea syndrome by increasing the number of apneas and hypopneas (<u>Ebben *et al*</u>, <u>2008</u>; <u>Holmes *et al*, 2003</u>; <u>Marzec *et al*, 2003; <u>Papacostas *et al*, 2007</u>). VNS should be used cautiously in patients with sleep apnea, or be supplemented with continuous positive airway pressure.</u>

Research Uses

Owing to the cost and invasive nature of VNS, there have been no human studies in healthy adults. Recently, some have proposed transcutaneous VNS (Huston et al, 2007; Kraus et al, 2007). However, studies in patients with epilepsy or depression implanted with VNS have shown that VNS causes discrete changes in limbic structures including the cingulate gyrus, the hippocampus, and the insula (Chae et al, 2003; Henry, 2002; Henry et al, 1999). The specific network activated depends on the choice of the use parameters (Mu et al, 2004), suggesting that with more extensive knowledge, one could 'direct' the VNS signal within groups of patients or even individually (Lomarev et al, 2002b). Human studies using fMRI and PET techniques show that VNS induces neuronal activity changes within amygdala, hippocampus, and thalamus, all targets of the LC (Henry et al, 1998, 1999; Lomarev et al, 2002a; Mu et al, 2004). These regional changes evolve over time and vary with clinical response (Nahas et al, 2007). Additionally, VNS produces interesting improvements in cognition (Boon et al, 2006; Borghetti et al, 2007; Helmstaedter et al, 2001; Sackeim et al, 2001a; Smith et al, 2006), perhaps linked to its influence on central LC norepinephrine system. Improvements in verbal recognition memory (Clark et al, 1999) and enhanced working memory (Sackeim et al, 2001b) have also been reported. VNS also has effects on sleep and arousal states. VNS decreases daytime sleepiness in humans (Malow et al, 2001) and promotes increased attention and arousal in animals (Lockard et al, 1990). These findings suggest that VNS may be a potent modulator of cognition through influences on ascending arousal systems. CSF studies have found increases in serotonin and norepinephrine metabolites following VNS.

The animal studies to date have been more extensive, although progress in this area was slowed by the lack of small portable generators. Now that these are available for rats, VNS studies have shown the importance of the LC in the signal propagation (Krahl *et al*, 1998), and have also shown long-term changes in raphe firing, unlike serotonin acting medications (Biggio *et al*, 2009; Dorr and Debonnel, 2006; Manta *et al*, 2009).

Studies in rodents have examined the functional relationship between the vagus nerve and LC, in addition to the anatomical circuit connections summarized above. VNS induces expression of the immediate early gene c-fos in LC neurons (<u>Naritokuet al, 1995</u>). Several studies have linked the LC to the seizure-suppressant effects of vagal activity. Thus, lesions of the LC attenuate the anti-epileptic effects of VNS in the rat (<u>Krahl et al, 1998</u>). Anatomical targets of LC projections also show electrophysiological and neurochemical changes following VNS. Amygdala, hippocampal, and insular cortex neurons all show enhanced neuronal activity after VNS (<u>Radna and MacLean, 1981a</u>, <u>1981b</u>). Microdialysis studies in animals show that VNS potentiates NE release in both the amygdala (<u>Hassert et al, 2004</u>) and hippocampus (<u>Miyashita and Williams, 2003</u>). VNS also induces c-fos expression in each of these structures as well as other LC targets including the thalamus (<u>Naritoku et al, 1995</u>). Thus, anatomical findings show that the vagus and LC are connected through well-specified relay nuclei, and functional studies show that these circuits contribute to forebrain activity. These findings show that chemical or electrical stimulation of the vagus nerve alters LC activity and that of its forebrain targets suggesting that the therapeutic effects of VNS may involve the LC-noradrenergic system.

There is also intense recent work investigating the role that VNS might have on inflammation and the immune response (<u>Ottani *et al*</u>, 2009; <u>Pavlov</u>, 2008; <u>Van Der Zanden *et al*</u>, 2009).

Clinical Studies

The first self-contained devices were implanted in humans in 1988 in patients with intractable, medically unresponsive epilepsy. Results were positive in two large acute double-blind controlled studies of VNS in patients with treatment-resistant epilepsy (<u>Ben-Menachem *et al*</u>, 1994; <u>Handforth *et al*, 1998</u>). Low-dose stimulation (intensity, number of pulses per day) served as the control in comparison to high stimulation. In this difficult to treat population, seizure frequency decreased 28–31% in the high stimulation group compared to baseline whereas it dropped 11–15% in the low stimulation group.

Unfortunately, few patients are able to stop their anticonvulsant medications although many are able to reduce the number of daily medications. This is clinically important in childhood epilepsy as many children experience deleterious cognitive side effects from the anticonvulsants (Ferrie and Patel, 2009; Shahwanet al, 2008).

Long-term follow-up studies have shown that the time course to respond to VNS is gradual, with continued improvement up to 1 year and then stabilization of effect. There appears to be no tolerance to VNS. The patient with the longest exposure to VNS has had the system operating for 17 years. VNS has assumed a small but significant role in epilepsy practice for those patients who have tried and failed two anticonvulsants.

VNS became available for use in Europe in 1994 and was given an FDA indication for epilepsy in the United States in 1997.

In 1997 one of the authors (MSG), along with John Rush, Harold Sackeim, and later Lauren Marangell, began an initial pilot study of VNS for patients with treatment-resistant depression (TRD) (<u>Rush *et al*</u>, <u>2000</u>; <u>Sackeim *et al*, 2001c</u>). Several lines of evidence suggested that VNS might be helpful in patients with depression, including anecdotal reports of mood improvement in VNS implanted epilepsy patients and functional imaging studies showing that VNS increased activity in several regions of the brain thought to be involved with depression (<u>Henry *et al*</u>, 1998</u>). This open-label study with 59 patients with TRD showed good results—30% response rate and 15% remission rate at 10 weeks. Even more encouraging were the extended results (<u>Marangell *et al*</u>, 2002; <u>Nahas *et al*</u>, 2005</u>). Patients continued to improve long after the acute phase of the trial. Patients were clinically better at 1 year than they were at 3 months. This pattern is unusual in the treatment of depression, especially in a difficult to treat cohort with prior tolerance to antidepressants (<u>Rush *et al*</u>, 2006a, 2006b</u>). A recent European trial found slightly better results, but with the same side effects and time course of response (Schlaepfer *et al*, 2008).

A pivotal multicentered, randomized, double-blinded trial of VNS was not as encouraging. In this underpowered trial, active VNS failed to statistically separate from sham treatment. The response rates for the acute treatment of TRD were 15% for active treatment and 10% for sham treatment (<u>Rush *et al*</u>, 2005a).

A parallel but nonrandomized group was also studied and compared with those patients who received VNS in the pivotal trial above. Thus, one group received the addition of VNS and the other received 'treatment as usual' (<u>Rush *et al*</u>, 2005b). They were followed for 12 months during which time both groups received similar treatment (medications and ECT) except for the VNS difference. At the end point the response rates were significantly different: 27% for the VNS group and 13% for the treatment as usual group (<u>George *et al*</u>, 2005).

The FDA considered all these studies when evaluating VNS for depression. They were most impressed with the long-term enduring benefits for this difficult to treat population (<u>George *et al*</u>, 2005). In 2005, they approved VNS for patients with chronic or recurrent depression, either unipolar or bipolar, with a history of failing to respond to at least four antidepressant trials. As VNS is FDA approved for TRD in the absence of Class I evidence of efficacy, insurance companies have resisted reimbursing the implant. Thus, currently VNS is not making a large clinical impact for depression treatment and the field awaits a much-needed adequately powered randomized controlled trial (RCT), which unfortunately has not been started because of financial concerns on the part of the manufacturer.

It is disappointing that the overall response rate to VNS plus medications at 1 year is less than 50%, as it is costly and requires a surgical implantation. However, many studies are now showing that patients with TRD have poor outcomes to traditional medication treatment (Fekadu *et al*, 2009; Rhebergen *et al*, 2009; Rush *et al*, 2006a, 2006b; Ten Doesschate *et al*, 2009; Trivedi *et al*, 2006; Yiend *et al*, 2009). Attempts to predict who is more likely to respond to VNS have not been successful.

There are several other potential VNS clinical applications, reasoning from the known role of the vagus, including obesity (Roslin and Kurian, 2001), craving (Bodenlos *et al*, 2007), pain (Borckardt *et al*, 2006a, 2005), traumatic brain injury (Colombo *et al*, 2008; Neese *et al*, 2007; Ottani *et al*, 2009), and anxiety (George*et al*, 2008b). These small sample size trials all suggest potential efficacies in these domains but RCT are needed.

TRANSCRANIAL MAGNETIC STIMULATION

Description of Method

Transcranial magnetic stimulation involves inducing an electrical current within the brain using pulsating magnetic fields that are generated outside the brain near the scalp. The essential feature is using electricity to generate a rapidly changing magnetic field, which in turn produces electrical impulses in the brain. A typical TMS device produces a fairly powerful magnetic field (about 1.5–3 T), but only very briefly (milliseconds). TMS is not simply applying a static or constant magnetic field to the brain. By 1820 scientists had discovered that passing an electric current though a wire induces a magnetic field generates an electrical current (Faraday, 1965). Thus, a changing magnetic field can generate electrical current in nearby wires, nerves, or muscles. A static magnet will not generate a current. For most TMS applications, it likely is the electricity induced from the pulsating magnet, and not the magnetic field itself, which produces neurobiological effects.

In 1959, Kolin and his colleagues showed that a fluctuating magnetic field could stimulate a peripheral frog muscle in preparation (Kolin *et al*, 1959). However, it was not until 1985 that the modern era of TMS started. That year Anthony Baker in Sheffield, England described the use of a noninvasive magnetic device resembling modern TMS instruments (Barker *et al*, 1985). The device was slow to recharge and quick to overheat, but it was able to stimulate spinal cord roots, and also superficial human cortex.

TMS requires a unit to store and deliver a charge (called a capacitor), and an electromagnetic coil (typically round in the shape of a doughnut or two round coils side-by-side and connected in a figure eight) (see Figure 2). A system can be cumbersome (resembling a small refrigerator), although some have shown that the entire system could be made portable and weigh less than 20 lbs (Epstein, 2008; Huang *et al*, 2009). The devices are regulated by the FDA for general safety, and most machines have FDA approval for sale in the US. They are also then regulated with respect to the ability to advertise their therapeutic use in a particular disorder. In the United States a device manufactured by Neuronetics was approved by the FDA in 2008 for treating depression (O'Reardon *et al*, 2007).

Figure 2.



Transcranial magnetic stimulation (TMS). Current from the wall (a) is used to charge a bank of large capacitors (b). These capacitors send a pulsing electrical current to the coils (c) resting on the scalp (d). The powerful but brief electrical current in the coil creates a transient magnetic field, which passes unimpeded through the skin and skull and results in electrical impulses in neurons in superficial cortex under the coil (e).

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Early TMS devices only emitted a single, brief pulse. Modern devices can generate a rapid succession of pulses, called repetitive TMS (rTMS). These devices are used for behavioral research or clinical treatments and can discharge on and off for several minutes. For example, the typical treatment for depression is a 20–40 min session, 5 days a week for 4–6 weeks. To keep the patient still and the device correctly placed, the patient reclines in a chair and the device is held securely against their head while they are awake and alert without needing anesthesia.

The TMS coil generates a magnetic field impulse that can only reach the outer layers of the cortex (<u>Davey *et al*</u>, 2004). The main effect of the impulse only penetrates 2–3 cm below the device (<u>Roth *et al*</u>, 1994; <u>Rothwell *et al*, 1999</u>). However, a deep TMS device has been invented and is in early clinical trials for depression and several other indications (<u>Roth *et al*</u>, 2002, 2005).

When the TMS device produces a pulse over the motor cortex, descending fibers are activated and volleys of electrical impulses descend through connected fibers into the spinal cord and out to the peripheral nerve where it can ultimately cause a muscle to twitch. The minimum amount of energy needed to produce contraction of the thumb (abductor pollicis brevis) is called the motor threshold (MT) (<u>Fitzgerald *et al*</u>, 2006; Fox *et al*, 2006; Sacco and Thickbroom, 2009). As this is so easy to generate, and varies widely across individuals, the MT is used as a measure of general cortical excitability and most TMS studies (both research and clinical) report the TMS intensity as a function of individual MT (and not as an absolute physical value) (<u>Di Lazzaro *et al*</u>, 2008). Although this convention has helped in making TMS safer, it is severely insufficient, in that it is referenced only to each machine, and thus is not a universal number. Future work is focusing on more universal, constant, measures of the magnetic field delivered.

In general with TMS, a stronger, more intense pulse results in more activation of the CNS tissue, and a wider area of activation. The circumstance with frequency is more complex. In general, frequencies of less than 1 per second (<1 Hz) are inhibitory (<u>Hoffman and Cavus, 2002</u>). This may be because low-frequency TMS more selectively stimulates the inhibitory GABA neurons, or this frequency is LTD like. Conversely, higher frequency stimulation is behaviorally excitatory (<u>Ziemannet al, 2008</u>). However, high-frequency TMS over some brain regions can temporarily block or knockout the function of that part of the brain (<u>Epstein et al</u>, 1996; Pascual-Leone et al, 1991).

A handheld device is being developed and studied as a treatment to interrupt migraine headaches (Neuralieve). The device delivers a single large pulse. When the patient experiences the aura phase of an impending headache they hold the device to the back of their head and direct the pulse toward the occipital cortex (Ambrosini and Schoenen, 2003; Clarke *et al*, 2006).

Putative Mechanisms of Action

TMS can produce different brain effects depending on the brain region being stimulated, the frequency of stimulation, the use parameters (intensity, frequency, duty train), and whether the brain region is engaged or 'resting'. Thus, it is difficult to review a single 'mechanism of action' for TMS. However, in general, a single pulse of TMS over a cortical region, such as the motor cortex, causes large neurons to depolarize. That is, the powerful transient magnetic field induces current to flow in neurons in superficial cortex (induced current). Both modeling and simple testing have shown that the fibers that are most likely to depolarize are those that are perpendicular to the coil, and are bending within the gyrus (Amassian et al, 1992, ; Lisanby et al, 1998a, 1998b). Some lower TMS intensities do not cause large neuron depolarization, but can still affect resting membrane potentials and thus alter brain activity and behavior. The most striking positive phenomena that TMS can produce are motor twitches (thumb, hand, arm, or leg movement) when applied over motor cortex, or 'phosphenes' when TMS is placed over the occiput. To date

TMS cannot produce acute memories, thoughts, or sensations or percepts apart from the scalp sensation of the coil.

rTMS can produce measurable effects lasting for minutes to hours after the train. In general, rTMS at frequencies greater than 1 Hz are excitatory, and less than 1 Hz inhibitory. One particular TMS sequence builds directly from the neurobiological studies of LTP and LTD, and is called theta burst as it has short bursts of TMS at theta frequencies (<u>Di Lazzaro *et al*</u>, 2005; <u>Stagg *et al*</u>, 2009).

TMS over some cortical regions can produce a transient disruption of behavior. This is most striking when the coil is placed over Broca's area and one can produce a transient expressive aphasia. Much interest involves whether TMS can produce short-term or even longer-term changes in plasticity (Ziemann *et al.*, 2008). Simple studies in motor and visual systems clearly indicate the potential for this approach (Miniussi *et al.*, 2008), which is now being applied in studies of poststroke recovery and other forms of rehabilitation (Hummel *et al.*, 2008; Pape*et al.*, 2009).

Coupling TMS with electrophysiological measures allows one to use TMS as a measure of motor cortex excitability, and then measure how behavior, medications, or other interventions might change excitability. Several groups are using this TMS excitability measurement technique to investigate new CNS-active compounds (Li *et al*, 2009, 2004; Paulus *et al*, 2008; Ziemann *et al*, 2008).

Coupling TMS with imaging (PET, SPECT, fMRI, or BOLD fMRI) allows one to directly stimulate circuits and then image the resultant changes (<u>George *et al*</u>, 2007; <u>Siebner *et al*, 2009</u>). With respect to the neuropsychiatric uses of TMS for depression or pain, at a molecular level TMS is known to have similar effects as those seen with ECT, for example, increased monoamine turnover, increased Brain-Derived Neurotrophic Factor, and normalization of the hypothalamic–pituitary–adrenal axis.

The initial use of daily prefrontal TMS to treat depression was based on the theory that clinical depression involved an imbalanced relationship between prefrontal cortex and limbic regions involved in mood regulation (insula, cingulate gyrus, amygdala, and hippocampus)(<u>George *et al*</u>, 1994</u>). There is only limited direct support that this is occurring, although recent work by Maier and colleagues directly supports the causal role of medial prefrontal cortex in mitigating and reversing chronic learned helplessness. Stimulatory fibers from PFCx are critical in this model (<u>Baratta *et al*</u>, 2007; <u>Christianson *et al*, 2008a, 2008b;Hutchinson *et al*, 2008).</u>

Safety

In general, TMS is regarded as safe and without enduring side effects. There have been no reported lasting neurologic, cognitive, or cardiovascular sequelae. However, TMS can alter brain function and is a relatively new technology so vigilance is required. The interested reader should read the results from an earlier international conference on TMS safety (<u>Wassermann, 1997</u>). An update has been drafted following another international meeting and should be available within the next 6 months.

Inducing a seizure is the primary safety concern with TMS. There have been less than 20 reported seizures induced with TMS, with a sample size of several thousand. The risk is less than one half of 1%. Most of these patients were healthy volunteers without a history of epilepsy. Fortunately, there are no reports that the individuals affected experienced recurrence. In addition, all of the seizures occurred during TMS administration when the patient was sitting down and near an investigator. In addition, all of the seizures were self-limited without needing medications or other interventions. Published safety tables concerning the proper intensity, frequency, and number of stimuli help minimize the numbers of seizures(<u>Wassermann, 1997</u>). Of the reported cases the majority were receiving TMS to the motor cortex—the most epileptogenic region of the cortex. Additionally, most (but not all) were receiving trains of stimulation outside of suggested limits. These cases suggest that TMS induced seizures will remain a small but significant adverse event even in patients without histories of seizures and even when TMS is used within suggested guidelines.

Studies in rabbits as well as some human studies suggest that TMS can cause hearing loss and subjects, patients, and operators should wear earplugs (<u>Counteret al, 1990</u>; <u>Loo et al, 2001</u>). One patient

reported a temporary hearing loss after TMS. In light of this an extensive study of auditory threshold was conducted before and after 4 weeks of TMS in over 300 patients. No changes were found. However, patients should wear earplugs when receiving TMS.

Headaches are the most common complaint after TMS, however, there was no difference in headache frequency between sham and control in a recent large trial (<u>O'Reardon *et al*</u>, 2007). Repeated analysis of neurocognitive functioning of TMS patients has not found any enduring negative effects from the procedure (<u>Avery*et al*</u>, 2008; <u>Little *et al*</u>, 2000). After a session, patients or subjects are able to drive home and return to work.

Research Uses

Space does not permit a thorough overview of TMS research uses, other than to highlight the active areas. TMS can be used as a measure of cortical excitability, and has been used to investigate medication effects, emotional states, plasticity in learning and stroke recovery, sleep (Massimini *et al*, 2007; Tononi and Koch, 2008), and in a host of disease states. TMS can be combined with brain imaging to directly stimulate circuits and image the resultant changes (see Figure 3). When precisely applied over critical brain regions, TMS can help causally determine whether a brain region is involved in a behavior, and how information flows through the brain during a task (Figure 4). There is much excitement, but little hard evidence, that TMS might be used to actually augment task performance, memory formation, or recovery from injury.

Figure 3.



Brain stimulation and imaging. The combination of brain imaging with brain stimulation allows for more direct examination of the role of circuit activity in brain behavior relationships. Historically most brain imaging has been relatively passive, and changes in a circuit occur along with a behavior, but causality is not known. By combining actual stimulation with imaging one can move a step closer to causal statements, as well as prepare the stage for potential clinical translation and therapeutic uses of brain stimulation approaches. In general, one can image simultaneously with stimulation (a), or one can use the brain imaging result (structural or functional or some combination) to guide the placement of the brain stimulation (in this case TMS) (b). Finally, one can stimulate a region with TMS or tDCS, produce brain changes, and then use brain imaging to examine changes in circuit behavior (c). (Reprinted with permission from Elsevier and adapted from Siebner *et al* (2009)).

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Figure 4.



State-dependent interregional interactions evoked by transcranial magnetic stimulation (TMS) interleaved with fMRI. Some groups can actually use TMS within an MRI scanner (Bohning *et al*, 1998). These images show the (a) main effect of left hand grip, irrespective of TMS stimulation intensity. This illustrates how one can obtain blood-oxygenation-level-dependent (BOLD) activation maps during concurrent application of TMS pulses (five pulses, 11 Hz) inside a magnetic resonance image (MRI) scanner. (b) Task-state-dependent effects of TMS on causal interactions in the human motor system. At rest, TMS applied to the left dorsal premotor cortex (PMd)

increased activity in contralateral PMd and primary motor cortex (M1) at high stimulation intensity (110% of resting motor threshold), compared with stimulation at a lower control intensity (70% active motor threshold). In contrast, this effect was reversed during a simple motor task that activated right PMd and M1. Now high-intensity stimulation increased task-related activity, compared with lower intensity stimulation. The results show how TMS can causally affect activity in contralateral regions, and that these influences are dependent on the activation state of these regions (adapted from <u>Bestmann *et al*(2003)</u> and reprinted with permission from Elsevier and <u>Siebner *et al*(2009</u>).

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Clinical Studies

Largely because of its noninvasiveness, TMS has been investigated in almost all neuropsychiatric conditions. Until only recently, there has not been a TMS industry to promote or perform this work and thus much of the clinical work has been single site and nonindustry funded, with relatively small sample sizes.

Depression has been the most widely studied condition with TMS. Three initial studies from Europe used TMS over the vertex as a potential antidepressant (Grisaru *et al.*, 1994; Hoflich *et al.*, 1993; Kolbinger *et al.*, 1995). In the US, George, Wassermann, and Post performed initial safety studies in healthy controls, an open study, and then a double-blind controlled trial of TMS for 2 weeks (George*et al.*, 1997, 1995, 1996). This work has now dramatically grown, but without much change in many of the initial treatment choices (coil location, frequency, dosing). There have now been several meta-analyses of the procedure (Ridding and Rothwell, 2007). A recent meta-analysis of rTMS for depression examined 25 published sham-controlled studies (Mitchell and Loo, 2006). The authors concluded that left prefrontal TMS provided statistical superiority over sham treatment for patients with depression. However, they concluded that the clinical benefits are marginal in the majority of reports and there is still considerable uncertainty concerning the optimal stimulation parameters. Two recent positive meta-analyses suggest that the overall effect size with TMS in major depression is at least as good as that of standard pharmacotherapy (Lam *et al.*, 2008;Schutter, 2008). Those clinical features that appear to be associated with greater response include younger age, lack of refractoriness to antidepressants, and no psychotic features (<u>Avery *et al.*, 2008</u>).

The largest multisite trial to date, which resulted in FDA approval, was by Neuronetics. They sponsored a double-blind, multisite study of 301 medication-free patients with major depression. Patients were randomized to active TMS or sham treatment, which they received for 4–6 weeks (<u>O'Reardon *et al*</u>,

<u>2007</u>). There was some controversy about the results of the trial. Before conducting the experiment, the company chose a continuous variable, the change from baseline on the Montgomery–Asberg Depression Rating Scale (MADRS), as the primary outcome measure (and did not tell investigators in the field) while using the Hamilton Rating Scale as the entry criteria. Unfortunately, at 6 weeks the continuous measured MADRS change from baseline for the active treatment group was not guite statistically different from the control group: p=0.058. The Hamilton Depression Rating Scale scores, considered secondary outcome measures, were indeed superior for those in the active treatment group. The company argued, successfully for the publication, that they should be able to exclude six subjects with entry MADRS scores that were very low and could not reflect clinical improvement. Thus, the manuscript was published as a positive trial but the FDA initially rejected the application, and only agreed for approval after reviewing response data on subgroups. As there was such a large effect seen in those who were less treatment resistant, the FDA labeling is for the treatment of MDD in adult patients who have failed to achieve satisfactory improvement from one prior antidepressant treatment at or above the minimal effective dose and duration in the current episode. Note that in clinical practice, only about one in four treatment trials meets criteria for minimal dose and duration, so this translates in a clinical practice to patients with a moderate level of treatment resistance (Dewet al, 2005; Joo et al, 2005; Oquendo et al, 2003).

These mixed results reflect the current status of TMS for depression. Most agree that daily left prefrontal TMS for several weeks has antidepressant effects and is safe and well tolerated. It will likely be an ideal treatment for some patients. However, the efficacy data in trials to date are not as robust as some would like and many await the results of larger ongoing trials and better understanding of the mechanisms of action. For example, a large European trial failed to find a statistically significant difference, but likely used

an active sham condition as well as examined TMS as an augmentation rather than stand-alone treatment (<u>Herwiget al, 2007</u>). The NIH has funded a large multisite trial in depression with results due in late 2009 and the VA has launched a large cooperative study of daily left prefrontal TMS in depressed veterans.

One recent development in terms of TMS positioning has highlighted that better understanding of the TMS methods used will likely boost clinical antidepressant efficacy. The early NIMH studies used a rough measurement technique known as the 5-cm rule to place the TMS coil roughly over the prefrontal cortex (George et al, 1997, 1995, 1996). As the location of the motor strip varies between individuals, and skull size (hat size) also varies, this simple rule results in a large variation of actual location on scalp. It became obvious that this was an insufficient technique, but was nevertheless used in most trials, including the one for FDA approval (Herwig et al, 2001). One study suggested that the 5-cm rule resulted in 30% of patients being treated over supplementary motor area rather than prefrontal cortex. Two retrospective analyses of clinical trials in which brain imaging was performed to document the coil location have independently confirmed that a coil position that is anterior and lateral is associated with a better clinical response to active but not sham TMS (Herbsman et al, 2009). An Australian group has performed a RCT and a more anterior and lateral location did indeed produce superior antidepressant response (Fitzgerald *et al*, 2009). These findings suggest that the TMS effect is not nonspecific, and that the location of the coil clearly matters, even within broad boundaries of a specific lobe. It is not clear whether individualized location will be needed or used, or whether general algorithms will suffice for most patients.

Auditory hallucinations are part of the positive symptoms of schizophrenia. These types of hallucinations are believed to result from aberrant activation of the language perception area at the junction of the left temporal and parietal cortices (<u>Higgins and George, 2007</u>). Low-frequency TMS has been used to potentially inhibit this area in patients with schizophrenia and provide relief from auditory hallucinations. A recent meta-analysis examined the efficacy of low-frequency TMS as a treatment of resistant auditory hallucinations in schizophrenia (<u>Aleman*et al*, 2007</u>). Ten sham-controlled studies have incorporated 212 patients. Their review concluded that TMS was effective in reducing auditory hallucinations. Unfortunately, TMS had no effect on other positive symptoms or the cognitive deficits of schizophrenia. Larger studies are needed to definitely establish the efficacy, tolerability, and utility of TMS for schizophrenia.

There have been four RCTs of using intermittent daily prefrontal TMS to treat negative symptoms in patients with schizophrenia. Only one of these studies was positive.

Tinnitus is a common, often disabling disorder, for which there is no adequate treatment. As many as 8% of adults over 50-years old suffer from tinnitus that can often be quite distressing. Recent functional imaging studies have identified increased activity in the auditory cortex in patients with tinnitus. Low-frequency TMS offers a possible mechanism to inhibit the overactive auditory cortex that may be producing tinnitus. Several small controlled trials from one research group in Germany have produced impressive results (Langguth *et al*, 2008). Larger, multicenter studies are needed to see if these positive effects can be replicated.

Numerous small controlled studies have evaluated the utility of TMS in patients with pain. Multiple sites have been tested including prefrontal cortex, motor cortex, and parietal cortex(<u>Andre-Obadia *et al*</u>, 2006; <u>Lefaucher *et al*, 2001; <u>Lefaucheur</u>, 2004; <u>Lefaucheur *et al*, 2001; <u>Pridmore and</u> <u>Oberoi, 2000</u>; <u>Rollnik *et al*, 2003</u>). In general, TMS provides effective pain relief in these different locations in diverse pain conditions. Unfortunately, the effect of TMS on pain only lasts for a short duration. Consequently, the utility of TMS as a practical treatment for chronic pain conditions has yet to be established.</u></u>

Recent studies suggest TMS may have some utility in managing acute pain. In two different studies of patients recovering from gastric by-pass surgery, 20 min of real or sham TMS was administered to the prefrontal cortex of every patient. Then their use of self-administered morphine was followed over the next 48 h. Those receiving real TMS used 40% less morphine in the next 24 h, with the majority of the reduction occurring in the first 8 h after TMS (<u>Borckardt *et al*</u>, 2008b, 2006b).

The handheld device, mentioned above, is being studied as a treatment for migraine headaches. Preliminary results have been encouraging. Larger studies are underway.

Following an ischemic event to the motor cortex, the brain attempts to reorganize the damaged networks. Indeed, the extent of reorganization correlates with the clinical recovery of motor function. TMS may accelerate the reorganization process and therefore enhance recovery (Hummel et al,

<u>2008</u>; <u>Miniussi et al, 2008</u>; <u>Pape et al, 2009</u>). It is unclear which types of TMS may be beneficial in stroke recovery. High-frequency TMS to the affected area may enhance reorganization. Alternatively, low-frequency TMS to the opposite, intact hemisphere is believed to reduce the interference from the nonstroke side. Some believe that too much input from the unaffected side of the brain impedes recovery. Reducing excitability with low-frequency TMS may enhance recovery.

Ridding and Rothwell recently reviewed the studies of TMS in stroke recovery. Although the total number of patients in controlled trials was only 87, the results were encouraging. Clearly, larger studies are needed, but it appears that TMS might be able to improve the natural healing process after a stroke (Kew *et al*, 1994; McKay *et al*, 2002; Ridding and Rothwell, 1995, 2007).

Theoretically low-frequency TMS could be used to treat cortical epilepsy. Early studies showed that TMS could reduce EEG epileptiform abnormalities. Initial case studies were positive. A controlled study of daily TMS by Theodore *et al* (2002) over the cortical site of seizures for 1 week found a statistically significant reduction in seizures. However, the authors concluded that TMS treatment was not clinically significant. More recently, in another controlled trial Cantello *et al*(2007) concluded that 'active' rTMS was no better than placebo for seizure reduction. Thus, the idea of using inhibitory doses of TMS to calm cortical targets is intriguing. However, the controlled trials to date have not been as successful.

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