Electrodiagnostic Studies of Nervous System Disorders: EEG, Evoked Potentials, and EMG

Michael J. Aminoff

ELECTROENCEPHALOGRAPHY

The electrical activity of the brain [the electroencephalogram (EEG)] is easily recorded from electrodes placed on the scalp. The potential difference between pairs of electrodes on the scalp (bipolar derivation) or between individual scalp electrodes and a relatively inactive common reference point (referential derivation) is amplified and displayed on a computer monitor, oscilloscope, or paper. The characteristics of the normal EEG depend on the patient’s age and level of arousal. The rhythmic activity normally recorded represents the postsynaptic potentials of vertically oriented pyramidal cells of the cerebral cortex and is characterized by its frequency. In normal awake adults lying quietly with the eyes closed, an 8- to 13-Hz alpha rhythm is seen posteriorly in the EEG, intermixed with a variable amount of generalized faster (beta) activity (>13 Hz); the alpha rhythm is attenuated when the eyes are opened (Fig. e45-1). During drowsiness, the alpha rhythm is also attenuated; with light sleep, slower activity in the theta (4–7 Hz) and delta (<4 Hz) ranges becomes more conspicuous.

Digital systems are now widely used for recording the EEG. They allow the EEG to be reconstructed and displayed with any desired format and manipulated for more detailed analysis, and also permit computerized techniques to be used to detect certain abnormalities. Activating procedures are generally undertaken while the EEG is recorded in an attempt to provoke abnormalities. Such procedures commonly include hyperventilation (for 3 or 4 min), photic stimulation, sleep, and sleep deprivation on the night prior to the recording.

Electroencephalography is relatively inexpensive and may aid clinical management in several different contexts.

THE EEG AND EPILEPSY

The EEG is most useful in evaluating patients with suspected epilepsy. The presence of electrographic seizure activity—i.e., of abnormal, repetitive, rhythmic activity having an abrupt onset and termination and a characteristic evolution—clearly establishes the diagnosis. The absence of such electrocerebral accompaniment does not exclude a seizure disorder, however, because there may be no change in the scalp-recorded EEG during certain focal seizures. With generalized tonic-clonic seizures, the EEG is always abnormal during the episode. It is often not possible to obtain an EEG during clinical events that may represent seizures, especially when such events occur unpredictably or infrequently. Continuous monitoring for prolonged periods in video-EEG telemetry units has made it easier to capture the electrocerebral accompaniments of such clinical episodes. Monitoring by these means is sometimes helpful in confirming that seizures are occurring, characterizing the nature of clinically equivocal episodes, and determining the frequency of epileptic events.

The EEG findings may also be helpful in the interictal period by showing certain abnormalities that are strongly supportive of a diagnosis of epilepsy. Such epileptiform activity consists of bursts of abnormal discharges containing spikes or sharp waves. The presence of epileptiform activity is not specific for epilepsy, but it

![Figure e45-1](image-url)

A. Normal EEG showing a posteriorly situated 9-Hz alpha rhythm that attenuates with eye opening. B. Abnormal EEG showing irregular diffuse slow activity in an obtunded patient with encephalitis. C. Irregular slow activity in the right central region, on a diffusely slowed background, in a patient with a right parietal glioma. D. Periodic complexes occurring once every second in a patient with Creutzfeldt-Jakob disease. Horizontal calibration: 1 s; vertical calibration: 200 μV in A; 300 μV in other panels. (From MJ Aminoff, ed: Electrodiagnosis in Clinical Neurology, 5th ed. New York, Churchill Livingstone, 2005.) In this and the following figure, electrode placements are indicated at the left of each panel and accord with the international 10:20 system. A, earlobe; C, central; F, frontal; P, parietal; T, temporal; O, occipital. Right-sided placements are indicated by even numbers, left-sided placements by odd numbers, and midline placements by Z.
has a much greater prevalence in epileptic patients than in normal individuals. However, even in an individual who is known to have epilepsy, the initial routine interictal EEG may be normal up to 60% of the time. Thus, the EEG cannot establish the diagnosis of epilepsy in many cases.

The EEG findings have been used in classifying seizure disorders and selecting appropriate anticonvulsant medication for individual patients (Fig. e45-2). The episodic generalized spike-wave activity that occurs during and between seizures in patients with typical absence epilepsy contrasts with focal interictal epileptiform discharges or ictal patterns found in patients with focal seizures. These latter seizures may have no correlates in the scalp-recorded EEG or may be associated with abnormal rhythmic activity of variable frequency, a localized or generalized distribution, and a stereotyped pattern that varies with the patient. Focal or lateralized epileptogenic lesions are important to recognize, especially if surgical treatment is contemplated. Intensive long-term monitoring of clinical behavior and the EEG is required for operative candidates, however, and this generally also involves recording from intracranially placed electrodes (which may be subdural, extradural, or intracerebral in location).

The findings in the routine scalp-recorded EEG may indicate the prognosis of seizure disorders: In general, a normal EEG implies a better prognosis than otherwise, whereas an abnormal background or profuse epileptiform activity suggests a poor outlook. The EEG findings are not helpful in determining which patients with head injuries, stroke, or brain tumors will go on to develop seizures, because in such circumstances epileptiform activity is commonly encountered regardless of whether seizures occur. The EEG findings are sometimes used to determine whether anticonvulsant medication can be discontinued in epileptic patients who have been seizure-free for several years, but the findings provide only a general guide to prognosis. Further seizures may occur after withdrawal of anticonvulsant medication despite a normal EEG or, conversely, may not occur despite a continuing EEG abnormality. The decision to discontinue anticonvulsant medication is made on clinical grounds, and the EEG does not have a useful role in this context except for providing guidance when there is clinical ambiguity or the patient requires reassurance about a particular course of action.

The EEG has no role in the management of tonic-clonic status epilepticus except when there is clinical uncertainty whether seizures are continuing in a comatose patient. In patients treated by pentobarbital-induced coma for refractory status epilepticus, the EEG findings are useful in indicating the level of anesthesia and whether seizures are occurring. During status epilepticus, the EEG shows repeated electrographic seizures or continuous spike-wave discharges. In nonconvulsive status epilepticus, a disorder that may not be recognized unless an EEG is performed, the EEG may also show continuous spike-wave activity (“spike-wave stupor”) or, less commonly, repetitive electrographic seizures (focal status epilepticus).

### THE EEG AND COMA

In patients with an altered mental state or some degree of obtundation, the EEG tends to become slower as consciousness is depressed, regardless of the underlying cause (Fig. e45-1). Other findings may also be present and may suggest diagnostic possibilities, as when electrographic seizures are found or there is a focal abnormality indicating a structural lesion. The EEG generally slows in metabolic encephalopathies, and triphasic waves may be present. The findings do not permit differentiation of the underlying metabolic disturbance but help to exclude other encephalopathic processes by indicating the diffuse extent of cerebral dysfunction. The response of the EEG to external stimulation is helpful prognostically because electrocerebral responsiveness implies a lighter level of coma than a nonreactive EEG. Serial records provide a better guide to prognosis than a single record and supplement the clinical examination in following the course of events. As the depth of coma increases, the EEG becomes nonreactive and may show a burst-suppression pattern, with bursts of mixed-frequency activity separated by intervals of relative cerebral inactivity. In other instances there is a reduction in amplitude of the EEG until eventually activity cannot be detected. Such electrocerebral silence does not necessarily reflect irreversible brain damage, because it may occur in hypothermic patients or with drug overdose. The prognosis of electrocerebral silence, when recorded using an adequate technique, depends upon the clinical context.

**Figure e45-2.** Electrographic seizures. A. Onset of a tonic seizure showing generalized repetitive sharp activity with synchronous onset over both hemispheres. B. Burst of repetitive spikes occurring with sudden onset in the right temporal region during a clinical spell characterized by transient impairment of external awareness. C. Generalized 3-Hz spike-wave activity occurring synchronously over both hemispheres during an absence (petit mal) attack. Horizontal calibration: 1 s; vertical calibration: 400 mV in A, 200 mV in B, and 750 mV in C. (From MJ Aminoff, ed: Electrodiagnosis in Clinical Neurology, 5th ed. New York, Churchill Livingstone, 2005.)
context in which it is found. In patients with severe cerebral anoxia, for example, electrocerebral silence in a technically satisfactory record implies that useful cognitive recovery will not occur.

In patients with clinically suspected brain death, an EEG, when recorded using appropriate technical standards, may be confirmatory by showing electrocerebral silence. However, complicating disorders that may produce a similar but reversible EEG appearance (e.g., hypothermia or drug intoxication) must be excluded. The presence of residual EEG activity in suspected brain death fails to confirm the diagnosis but does not exclude it. The EEG is usually normal in patients with locked-in syndrome and helps in distinguishing this disorder from the comatose state with which it is sometimes confused clinically.

The brief EEG obtained routinely in the laboratory often fails to distinguish between dementia and pseudodementia. The EEG is normal or diffusely slowed in most patients with dementias, the EEG is normal or diffusely slowed, and the EEG findings alone cannot indicate whether a patient is demented or distinguish between dementia and pseudodementia.

Continuous EEG monitoring

The brief EEG obtained routinely in the laboratory often fails to reveal abnormalities that are transient and infrequent. Continuous monitoring over 12 or 24 hours or longer may detect abnormalities or capture clinical events that would otherwise be missed. The EEG is often recorded continuously in critically ill patients to detect early changes in neurologic status, which is particularly useful when the clinical examination is limited. Continuous EEG recording in this context has been used to detect acute events such as nonconvulsive seizures or developing cerebral ischemia, to monitor cerebral function, and to provide early warning of the occurrence of acute events such as focal or diffuse slowing or the appearance of spike-and-wave activity.

Evoked potentials

The noninvasive recording of spinal or cerebral potentials elicited by stimulation of specific afferent pathways is an important means of monitoring the functional integrity of these pathways but does not indicate the pathologic basis of lesions involving them. Such evoked potentials (EPs) are so small compared to the background EEG activity that the responses to a number of stimuli have to be recorded and averaged with a computer in order to permit their recognition and definition. The background EEG activity, which has no fixed temporal relationship to the stimulus, is averaged out by this procedure.

Visual evoked potentials (VEPs) are elicited by monocular stimulation with a reversing checkerboard pattern and are recorded from the occipital region in the midline and on either side of the scalp. The component of major clinical importance is the so-called P100 response, a positive peak having a latency of approximately 100 ms. Its presence, latency, and symmetry over the two sides of the scalp are noted. Amplitude may also be measured, but changes in size are much less helpful for the recognition of pathology. VEPs are most useful in detecting dysfunction of the visual pathways anterior to the optic chiasm. In patients with acute severe optic neuritis, the P100 is frequently lost or grossly attenuated; as clinical recovery occurs and visual acuity improves, the P100 is restored but with an increased latency that generally remains abnormally prolonged indefinitely. The VEP findings are therefore helpful in indicating previous or subclinical optic neuritis. They may also be abnormal with ocular abnormalities and with other causes of optic nerve disease, such as ischemia or compression by a tumor. Normal VEPs may be elicited by flash stimuli in patients with cortical blindness. Routine VEPs record a mass response over a relatively large cortical area and thus may be insensitive to localized waveform abnormalities. A newer technique, multifocal VEP, measures responses from 120 individual sectors within each affected eye, and thus is likely to be more sensitive than routine VEP.

Brainstem auditory evoked potentials (BAEPs) are elicited by monaural stimulation with repetitive clicks and are recorded between the vertex and the mastoid process or earlobe. A series of potentials, designated by roman numerals, occurs in the first 10 ms after the stimulus and represents in part the sequential activation of different structures in the pathway between the auditory nerve (wave I) and the inferior colliculus (wave V) in the midbrain. The presence, latency, and interpeak latency of the first five positive potentials recorded at the vertex are evaluated. The findings are helpful in screening for acoustic neuromas, detecting brainstem pathology, and evaluating comatose patients. The BAEPs are normal in coma due to metabolic/toxic disorders or bihemispheric disease but abnormal in the presence of brainstem pathology.

Somatosensory evoked potentials (SSEPs) are recorded over the scalp and spine in response to electrical stimulation of a peripheral (mixed or cutaneous) nerve. The configuration, polarity, and latency of the responses depend on the nerve that is stimulated and on the recording arrangements. SSEPs are used to evaluate proximal (otherwise inaccessible) portions of the peripheral nervous system and the integrity of the central somatosensory pathways.

Clinical utility of EPs

EP studies may detect and localize lesions in afferent pathways in the central nervous system (CNS). They have been used particularly
to investigate patients with suspected multiple sclerosis (MS), the
diagnosis of which requires the recognition of lesions involving
several different regions of the central white matter. In patients with
clinical evidence of only one lesion, the electrophysiologic recogni-
tion of abnormalities in other sites helps to suggest or support the
diagnosis but does not establish it unequivocally. Multimodality
EP abnormalities are not specific for MS; they may occur in AIDS,
Lyme disease, systemic lupus erythematosus, neurosyphilis, spino-
cerebellar degenerations, familial spastic paraplegia, and deficiency
of vitamin E or B₁₂, among other disorders. The diagnostic utility
of the electrophysiologic findings therefore depends on the circum-
stances in which they are found. Abnormalities may aid in the local-
ization of lesions to broad areas of the CNS, but attempts at precise
localization on electrophysiologic grounds are misleading because
the generators of many components of the EP are unknown.

The EP findings are sometimes of prognostic relevance. Bilateral
loss of SSEP components that are generated in the cerebral cortex
implies that cognition may not be regained in posttraumatic or
postanoxic coma, and EP studies may also be useful in evaluating
patients with suspected brain death. In patients who are comatose
for uncertain reasons, preserved BAEPs suggest either a metaboli-
toxic etiology or bitemporal hypoxic disease. In patients with spinal
cord injuries, SSEPs have been used to indicate the completeness
of the lesion. The presence or early return of a cortically generated
response to stimulation of a nerve below the injured segment of the
cord indicates an incomplete lesion and thus a better prognosis for
functional recovery than otherwise. In surgery, intraoperative EP
monitoring of neural structures placed at risk by the procedure may
permit the early recognition of dysfunction and thereby permit a
neurologic complication to be averted or minimized.

Visual and auditory acuity may be determined using EP tech-
niques in patients whose age or mental state precludes traditional
ophthalmologic or audiologic examinations.

■ COGNITIVE EVOKED POTENTIALS

Certain EP components depend on the mental attention of the
subject and the setting in which the stimulus occurs, rather than
simply on the physical characteristics of the stimulus. Such “event-
related” potentials (ERPs) or “endogenous” potentials are related
in some manner to the cognitive aspects of distinguishing an
infrequently occurring target stimulus from other stimuli occurring
more frequently. For clinical purposes, attention has been directed
particularly at the so-called P3 component of the ERP, which is
also designated the P300 component because of its positive polarity
and latency of approximately 300–400 ms after onset of an audi-
tory target stimulus. The P3 component is prolonged in latency
in many patients with dementia, whereas it is generally normal in
patients with depression or other psychiatric disorders that might
be mistaken for dementia. ERP’s are, therefore, sometimes helpful in
making this distinction when there is clinical uncertainty, although
a response of normal latency does not exclude dementia.

■ MOTOR EVOKED POTENTIALS

The electrical potentials recorded from muscle or the spinal cord
following stimulation of the motor cortex or central motor path-
ways are referred to as motor evoked potentials. For clinical purposes
such responses are recorded most often as the compound muscle
action potentials elicited by transcutaneous magnetic stimulation
of the motor cortex. A strong but brief magnetic field is produced
by passing a current through a coil, and this induces stimulating
currents in the subjacent neural tissue. The procedure is painless
and apparently safe. Abnormalities have been described in several
neurologic disorders with clinical or subclinical involvement of
central motor pathways, including MS and motor neuron disease.

In addition to a possible role in the diagnosis of neurologic dis-
orders or in evaluating the extent of pathologic involvement, the
technique provides information of prognostic relevance (e.g., in
suggesting the likelihood of recovery of motor function after stroke)
and provides a means of monitoring intraoperatively the functional
integrity of central motor tracts. Nevertheless, it is not used widely
for clinical purposes.

■ ELECTROPHYSIOLOGIC STUDIES OF MUSCLE AND NERVE

The motor unit is the basic element subserving motor function. It is
defined as an anterior horn cell, its axon and neuromuscular jun-
tions, and all the muscle fibers innervated by the axon. The number
of motor units in a muscle ranges from approximately 10 in the
extraocular muscles to several thousand in the large muscles of the
legs. There is considerable variation in the average number of mus-
cle fibers within the motor units of an individual muscle, i.e., in,
the innervation ratio of different muscles. Thus the innervation ratio
<25 in the human external rectus or platysma muscle and between
1600 and 1700 in the medial head of the gastrocnemius muscle. The
muscle fibers of individual motor units are divided into two general
types by distinctive contractile properties, histochemical stains, and
characteristic responses to fatigue. Within each motor unit, all of
the muscle fibers are of the same type.

■ ELECTROMYOGRAPHY

The pattern of electrical activity in muscle [i.e., the electromyogram
(EMG)], both at rest and during activity, may be recorded from a
needle electrode inserted into the muscle. The nature and pattern
of abnormalities relate to disorders at different levels of the motor
unit. Relaxed muscle normally is electrically silent except in the end
plate region, but abnormal spontaneous activity (Fig. e45-3) occurs
in various neuromuscular disorders, especially those associated with
denervation or inflammatory changes in affected muscle. Fibrillation
potentials and positive sharp waves (which reflect muscle fiber
irritability) and complex repetitive discharges are most often—but
not always—found in denervated muscle and may also occur after
muscle injury and in certain myopathic disorders, especially
inflammatory disorders such as polymyositis. After an acute
neuropathic lesion, they are found earlier in proximal rather than distal
muscles and sometimes do not develop distally in the extremities for

Figure e45-3 Activity recorded during EMG. A. Spontaneous fibrilla-
tion potentials and positive sharp waves. B. Complex repetitive discharges
recorded in partially denervated muscle at rest. C. Normal triphasic motor
unit action potential. D. Small, short-duration, polyphasic motor unit
action potential such as is commonly encountered in myopathic disorders.
E. Long-duration polyphasic motor unit action potential such as may be seen
in neuropathic disorders.
4–6 weeks; once present, they may persist indefinitely unless reinervation occurs or the muscle degenerates so completely that no viable tissue remains. Fasciculation potentials (which reflect the spontaneous activity of individual motor units) are characteristic of slowly progressive neuropathic disorders, especially those with degeneration of anterior horn cells (such as amyotrophic lateral sclerosis). Myotonic discharges—high-frequency discharges of potentials derived from single muscle fibers that wax and wane in amplitude and frequency—are the signature of myotonic disorders such as myotonic dystrophy or myotonia congenita but occur occasionally in polymyositis or other, rarer, disorders.

Slight voluntary contraction of a muscle leads to activation of a small number of motor units. The potentials generated by any muscle fibers of these units that are within the pickup range of the needle electrode will be recorded (Fig. e45-3). The parameters of normal motor unit action potentials depend on the muscle under study and age of the patient, but their duration is normally between 5 and 15 ms, amplitude is between 200 μV and 2 mV, and most are bi- or triphasic. The number of units activated depends on the degree of voluntary activity. An increase in muscle contraction is associated with an increase in the number of motor units that are activated (recruited) and in the frequency with which they discharge. With a full contraction, so many motor units are normally activated that individual motor unit action potentials can no longer be distinguished, and a complete interference pattern is said to have been produced.

The incidence of small, short-duration, polyphasic motor unit action potentials (i.e., having more than four phases) is usually increased in myopathic muscle, and an excessive number of units is activated for a specified degree of voluntary activity. By contrast, the loss of motor units that occurs in neuropathic disorders leads to a reduction in number of units activated during a maximal contraction and an increase in their firing rate, i.e., there is an incomplete or reduced interference pattern. The configuration and dimensions of the potentials may also be abnormal, depending on the duration of the neuropathic process and on whether reinnervation has occurred. The surviving motor units are initially normal in configuration but, as reinnervation occurs, they increase in amplitude and duration and become polyphasic (Fig. e45-3).

Action potentials from the same motor unit sometimes fire with a consistent temporal relationship to each other, so that double, triple, or multiple discharges are recorded, especially in tetany, hemifacial spasm, or myokymia.

Electrical silence characterizes the involuntary, sustained muscle contraction that occurs in phosphorylase deficiency, which is designated a contracture.

EMG enables disorders of the motor units to be detected and characterized as either neurogenic or myopathic. In neurogenic disorders, the pattern of affected muscles may localize the lesion to the anterior horn cells or to a specific site as the axons traverse a nerve root, limb plexus, and peripheral nerve to their terminal arborizations. The findings do not enable a specific etiologic diagnosis to be made, however, except in conjunction with the clinical findings and results of other laboratory studies.

The findings may provide a guide to the severity of an acute disorder of a peripheral or cranial nerve (by indicating whether denervation has occurred and the completeness of the lesion) and whether the pathologic process is active or progressive in chronic or degenerative disorders such as amyotrophic lateral sclerosis. Such information is important for prognostic purposes.

Various quantitative EMG approaches have been developed. The most common is to determine the mean duration and amplitude of 20 motor unit action potentials using a standardized technique. The technique of macro-EMG provides information about the number and size of muscle fibers in a larger volume of the motor unit territory and has also been used to estimate the number of motor units in a muscle. Scanning EMG is a computer-based technique that has been used to study the topography of motor unit action potentials and, in particular, the spatial and temporal distribution of activity in individual units. The technique of single-fiber EMG is discussed separately below.

**NERVE CONDUCTION STUDIES**

Recording of the electrical response of a muscle to stimulation of its motor nerve at two or more points along its course (Fig. e45-4) permits conduction velocity to be determined in the fastest-conducting motor fibers between the points of stimulation. The latency and amplitude of the electrical response of muscle (i.e., of the compound muscle action potential) to stimulation of its motor nerve at a distal site are also compared with values defined in normal subjects. Sensory nerve conduction studies are performed by determining the conduction velocity and amplitude of action potentials in sensory fibers when these fibers are stimulated at one point and the responses are recorded at another point along the course of the nerve. In adults, conduction velocity in the arms is normally between 50 and 70 m/s, and in the legs is between 40 and 60 m/s.

Nerve conduction studies complement the EMG examination, enabling the presence and extent of peripheral nerve pathology to be determined. They are particularly helpful in determining whether sensory symptoms are arising from pathology proximal or distal to the dorsal root ganglia (in the former instance, peripheral sensory conduction studies will be normal) and whether neuromuscular dysfunction relates to peripheral nerve disease. In patients with a mononeuropathy, they are invaluable as a means of localizing...
afferent fibers constitute the afferent arc and alpha motor axons of the nerve and represents a monosynaptic reflex in which spindle (Ia) nerve terminals). Such antidromic impulses cause a few of the afferent fibers to discharge, producing a small motor response that occurs considerably later than the direct response elicited by nerve stimulation. The F wave so elicited is sometimes abnormal (absent or delayed) with proximal pathology of the peripheral nervous system, such as a radiculopathy, and may therefore be helpful in detecting abnormalities when conventional nerve conduction studies are normal. In general, however, the clinical utility of F-wave studies has been disappointing, except perhaps in Guillain-Barré syndrome, where they are often absent or delayed.

**H-REFLEX STUDIES**

The H reflex is easily recorded only from the soleus muscle (S1) in normal adults. It is elicited by low-intensity stimulation of the tibial nerve and represents a monosynaptic reflex in which spindle (Ia) afferent fibers constitute the afferent arc and alpha motor axons of the efferent pathway. The H reflexes are often absent bilaterally in elderly patients or with polyneuropathies and may be lost unilaterally in S1 radiculopathies.

**MUSCLE RESPONSE TO REPETITIVE NERVE STIMULATION**

The size of the electrical response of a muscle to supramaximal electrical stimulation of its motor nerve relates to the number of muscle fibers that are activated. Neuromuscular transmission can be tested by several different protocols, but the most helpful is to record with surface electrodes the electrical response of a muscle to supramaximal stimulation of its motor nerve by repetitive (2–3 Hz) shocks delivered before and at selected intervals after a maximal voluntary contraction.

There is normally little or no change in size of the compound muscle action potential following repetitive stimulation of a motor nerve at 2–3 Hz with stimuli delivered at intervals after voluntary contraction of the muscle for about 20–30 s, even though preceding activity in the junctional region influences the release of acetylcholine and thus the size of the end-plate potentials elicited by a test stimulus. This is because more acetylcholine is normally released than is required to bring the motor end-plate potentials to the threshold for generating muscle fiber action potentials. In disorders of neuromuscular transmission this safety factor is reduced. Thus in myasthenia gravis, repetitive stimulation, particularly at a rate of between 2 and 5 Hz, may lead to a depression of neuromuscular transmission, with a decrement in size of the response recorded from affected muscles. Similarly, immediately after a period of maximal voluntary activity, single or repetitive stimuli of the motor nerve may elicit larger muscle responses than before, indicating that more muscle fibers are responding. This postactivation facilitation of neuromuscular transmission is followed by a longer-lasting period of depression, maximal between 2 and 4 min after the conditioning period and lasting for as long as 10 min or so, during which responses are reduced in size.

Decrementing responses to repetitive stimulation at 2–5 Hz are common in myasthenia gravis but may also occur in the congenital myasthenic syndromes. In Lambert Eaton myasthenic syndrome, in which there is defective release of acetylcholine at the neuromuscular junction, the compound muscle action potential elicited by a single stimulus is generally very small. With repetitive stimulation at rates of up to 10 Hz, the first few responses may decline in size, but subsequent responses increase. If faster rates of stimulation are used (20–50 Hz), the increment may be dramatic so that the amplitude of compound muscle action potentials eventually reaches a size that is several times larger than the initial response. In patients with botulism, the response to repetitive stimulation is similar to that in Lambert Eaton myasthenic syndrome, although the findings are somewhat more variable and not all muscles are affected.

**F-WAVE STUDIES**

Stimulation of a motor nerve causes impulses to travel antidromically (i.e., toward the spinal cord) as well as orthodromically (to the nerve terminals). Such antidromic impulses cause a few of the anterior horn cells to discharge, producing a small motor response that occurs considerably later than the direct response elicited by nerve stimulation. The F wave so elicited is sometimes abnormal (absent or delayed) with proximal pathology of the peripheral nervous system, such as a radiculopathy, and may therefore be helpful in detecting abnormalities when conventional nerve conduction studies are normal. In general, however, the clinical utility of F-wave studies has been disappointing, except perhaps in Guillain-Barré syndrome, where they are often absent or delayed.

**SINGLE-FIBER ELECTROMYOGRAPHY**

This technique is particularly helpful in detecting disorders of neuromuscular transmission. A special needle electrode is placed within a muscle and positioned to record action potentials from two muscle fibers belonging to the same motor unit. The time interval between the two potentials will vary in consecutive discharges; this is called the *neuromuscular jitter*. The jitter can be quantified as the mean difference between consecutive interpotential intervals and is normally between 10 and 50 μs. This value is increased when neuromuscular transmission is disturbed for any reason, and in some instances impulses in individual muscle fibers may fail to occur because of impulse blocking at the neuromuscular junction. Single-fiber EMG is more sensitive than repetitive nerve stimulation or determination of acetylcholine receptor antibody levels in diagnosing myasthenia gravis.
Single-fiber EMG can also be used to determine the mean fiber density of motor units (i.e., mean number of muscle fibers per motor unit within the recording area) and to estimate the number of motor units in a muscle, but this is of less immediate clinical relevance.

**BLINK REFLEXES**

Electrical or mechanical stimulation of the supraorbital nerve on one side leads to two separate reflex responses of the orbicularis oculi—an ipsilateral R1 response having a latency of approximately 10 ms and a bilateral R2 response with a latency in the order of 30 ms. The trigeminal and facial nerves constitute the afferent and efferent arcs of the reflex, respectively. Abnormalities of either nerve or intrinsic lesions of the medulla or pons may lead to uni- or bilateral loss of the response, and the findings may therefore be helpful in identifying or localizing such pathology.

**FURTHER READINGS**


