The Sensorimotor System, Part I: The Physiologic Basis of Functional Joint Stability

Bryan L. Riemann; Scott M. Lephart

University of Pittsburgh, Pittsburgh, PA

Objective: To define the nomenclature and physiologic mechanisms responsible for functional joint stability. Data Sources: Information was drawn from an extensive MEDLINE search of the scientific literature conducted in the areas of proprioception, neuromuscular control, and mechanisms of functional joint stability for the years 1970 through 1999. An emphasis was placed on defining pertinent nomenclature based on the original references. Data Synthesis: Afferent proprioceptive input is conveyed to all levels of the central nervous system. They serve fundamental roles in optimal motor control and sensorimotor control over functional joint stability.

Conclusions/Applications: Sensorimotor control over the dynamic restraints is a complex process that involves components traditionally associated with motor control. Recognizing and understanding the complexities involved will facilitate the continued development and institution of management strategies based on scientific rationales.

Key Words: proprioception, neuromuscular, motor control

The purpose of this 2-part series is to provide an overview of the current understanding surrounding peripheral afferent information acquisition and processing and levels of motor control as they relate to functional joint stability. We recognize that these papers focus heavily upon basic science research that, in many circumstances, lacks immediate clinical application. Our premise is to present the athletic training community with an introduction concerning how the dynamic restraints are activated and controlled by the motor control system of the body. Our goal is that these papers may initiate common understanding regarding the terminology and underlying physiology associated with proprioception and neuromuscular control. Ultimately, by establishing a baseline understanding about the sensorimotor system, clinical techniques can continue to be developed and applied with scientific rationale. Furthermore, through this understanding, clinicians can appreciate future developments and research directions focusing on the restoration of functional joint stability.

The purpose of this first paper is to introduce the sensorimotor motor system, the biological system that controls the contributions of the dynamic restraints for functional joint stability. A secondary purpose is to define the nomenclature pertaining to the mechanisms responsible for both the sensory and motor components of proprioception and neuromuscular control for the maintenance of functional joint stability.

PERTINENT TERMINOLOGY

Before examining the specialized components and physiologic intricacies of the sensorimotor system, we must begin our discussion by defining some broad terms used in the medical and physiologic literature. Homeostasis is defined as the dynamic process by which an organism maintains and controls its internal environment despite perturbations from external forces.

Because cells, tissues, and organs operating within the body can only function within narrow ranges of environmental conditions, maintaining homeostasis becomes the major driving force underlying many, if not all, physiologic functions of the body. The body is composed of many systems that operate automatically and unconscious. Systems are organized hierarchically, beginning at the cellular level, and contribute to bodily homeostasis in specific domains. In a healthy individual, the system’s homeostasis is usually maintained by 2 different control systems. Stimulation of a corrective response within the corresponding system after sensory detection is often considered feedback controls. In contrast, feedforward controls have been described as anticipatory actions occurring before the sensory detection of a homeostatic disruption.

Initiated feedback actions are largely shaped by previous experience with the detected stimulus. Somatosensory, visual, and vestibular input provides the information necessary for both forms of control during motor activities; however, the methods of information processing differ. Feedback control is characterized by a continual processing of afferent information, providing response control on a moment-to-moment basis. In contrast, afferent information during feedforward control is used intermittently until feedback controls are initiated.
Unfortunately, classifying an action as either feedback or feedforward is not as straightforward as their definitions suggest. In some circumstances, a combination of both feedforward and feedback control exists, such as during the maintenance of postural control. Additionally, consider the situation in which a subject watches a tester trigger a device that induces a joint perturbation. Many subjects will naturally "tense up" when they see the tester beginning to push the trigger before the perturbation. Whether the muscle activation before the perturbation reaching the joint is the result of feedforward or feedback control remains controversial. For this reason, the term feedforward control has been recommended to describe actions occurring upon the identification of the beginning, as well as the effects, of an impending event or stimulus. In contrast, feedback control should be used to describe actions occurring in response to the sensory detection of direct effects from the arrival of the event or stimulus to the system.

The actions occurring with both feedback and feedforward controls involve the hierarchic organization of a system, beginning at the cellular level and extending through both the tissue and organ levels. The action patterns used to restore homeostasis are defined as mechanisms. For example, the reflexive response is the mechanism the body uses to maintain or restore joint stability after an imposed joint perturbation. Within a given mechanism are multiple processes that ultimately lead to the achievement of the result. In the case of joint perturbation, the processes include mechanoreceptor stimulation, neural transmission, integration of the signals by the central nervous system (CNS), transmission of an efferent signal, muscle activation, and force production. By definition, for the purposes of this paper, assessing a mechanism refers to the cumulative outcome of the underlying processes. During many clinical and research evaluations, inferences about the integrity of mechanisms are made by measuring specific characteristics of the underlying processes. Onset latency of muscle activation, as measured through electromyography, is frequently assessed in joint perturbation.

One additional physiologic term requiring attention in a broad context before our specialized discussion is stability. Stability is defined as the state of remaining unchanged, even in the presence of forces that would normally change the state or condition. It has been further described as the property of returning to an initial state upon disruption. With respect to joints, based on the above definitions, we define stability as the state of a joint remaining or promptly returning to proper alignment through an equalization of forces.

THE SENSORIMOTOR SYSTEM

The sensorimotor system, a subcomponent of the comprehensive motor control system of the body, is extremely complex. The term sensorimotor system was adopted by the participants of the 1997 Foundation of Sports Medicine Education and Research workshop to describe the sensory, motor, and central integration and processing components involved in maintaining joint homeostasis during bodily movements (functional joint stability) (Figure 1). The components giving rise to functional joint stability must be flexible and adaptable because the required levels vary among both persons and tasks. The process of maintaining functional joint stability is accomplished through a complementary relationship between static and dynamic components. Ligaments, joint capsule, cartilage, friction, and the bony geometry within the articulation comprise the static (passive) components. Dynamic contributions arise from feedforward and feedback neuromotor control over the skeletal muscles crossing the joint. Underlying the effectiveness of the dynamic restraints are the biomechanical and physical characteristics of the joint. These characteristics include range of motion and muscle strength and endurance.

From these descriptions of static and dynamic stability components, it becomes apparent that the terms are not synonymous. Integrity of static stabilizers is measured through clinical joint stress testing (ligamentous laxity testing) and arthrometry, giving rise to the frequently used term clinical stability. Because of the complexity of the control over the dynamic restraints, measuring dynamic stability is more challenging. Currently, as described in a companion paper, we are only able to quantitatively measure certain characteristics of the dynamic stability mechanism.

PROPRIOCEPTION AND NEUROMUSCULAR CONTROL

Proprioception predominates as the most misused term within the sensorimotor system. It has been incorrectly used synonymously and interchangeably with kinesthesia, joint position sense, somatosensation, balance, and reflexive joint stability. In Sherrington’s original description of the “proprioceptive system,” proprioception was used to reference the afferent information arising from “proprioceptors” located in the “proprioceptive field.” The “proprioceptive field” was specifically defined as that area of the body “screened from the environment” by the surface cells, which contained recep-

Figure 1. The sensorimotor system incorporates all the afferent, efferent, and central integration and processing components involved in maintaining functional joint stability. Although visual and vestibular input contributes, the peripheral mechanoreceptors are the most important from a clinical orthopaedic perspective. The peripheral mechanoreceptors (pictured on the lower left) reside in the cutaneous, muscular, joint, and ligamentous tissues. Afferent pathways (dotted lines) convey input from the 3 levels of motor control and associated areas such as the cerebellum. Activation of motor neurons may occur in direct response to peripheral sensory input (reflexes) or from descending motor commands, both of which may be modulated or regulated by the muscle (gray lines). Efferent pathways from each of the motor control levels (solid lines) converge upon the alpha and gamma motor neurons located in the central aspects of the spinal cord. The contractions by the extraxial and intraxial muscle fibers cause new stimuli to be presented to the peripheral mechanoreceptors.
tors specially adapted for the changes occurring inside the organism independent of the “interoceptive field” (alimentary canal and viscera organs). In several of his writings, Sherrington declared proprioception as being used for the regulation of total posture (postural equilibrium) and segmental posture (joint stability), as well as initiating several conscious peripheral sensations (“muscle senses”). Although he considered vestibular information to be proprioceptive with respect to the head, Sherrington clearly delineated the functions of the labyrinth from those receptors in the periphery. According to Matthews, Sherrington described 4 submodalities of “muscle sense” in Schafer’s Textbook of Physiology: (1) posture, (2) passive movement, (3) active movement, and (4) resistance to movement. These submodality sensations correspond to the contemporary terms joint position sense (posture of segment), kinesthesia (active and passive), and the sense of resistance or heaviness. Thus, proprioception correctly describesafferent information arising from internal peripheral areas of the body that contribute to postural control, joint stability, and several conscious sensations.

In contrast to proprioception, the term somatosensory (or somatosensation) is more global and encompasses all of the mechanoreceptive, thermoreceptive, and pain information arising from the periphery. Conscious appreciation of somatosensory information leads to the sensations of pain, temperature, tactile (ie, touch, pressure, etc), and the conscious submodality proprioception sensations. Thus, as Figure 2 illustrates, conscious appreciation of proprioception is a subcomponent of somatosensation and, therefore, the terms should not be used interchangeably.

Although Sherrington’s definition of the proprioceptive field clearly excludes the receptors sensitive to the external environment (“exteroceptive field”), he did not imply that the receptors in each region function in total exclusion of one another. Rather, Sherrington recognized the interaction between receptors located in both regions of the body, referring to the relationship between the receptors in the exteroceptive and proprioceptive environments as “allied.” Specifically, with respect to conscious proprioception appreciation, this aspect of proprioception has undoubtedly led to much of the confusion surrounding the interpretation of conscious proprioceptive acuity in persons suspected of having diminished proprioceptive information arising from articular sources following orthopaedic injury. Care is required to differentiate between the sources of proprioception and the conscious sensations of proprioception because receptors located in the proprioceptive field may not be the only contributory sources. Depending upon the exact circumstances of a situation or task, sources contributing to conscious sensations of proprioception (ie, joint position sense) could potentially include the deeper receptors (ie, joint and muscle mechanoreceptors) typically associated with proprioception or the more superficial receptors that elicit tactile sensations, or both. Therefore, although the proprioception and tactile sensations are considered to be distinctly different sensory phenomena, similar sensory organs may contribute to each conscious sensation under particular conditions. A complete discussion of the sources contributing to conscious proprioception perception is presented in a later section of this paper.

Lastly, mechanoreceptors conveying proprioceptive information are often labeled as proprioceptors. However, in addition to mechanoreceptors located in Sherrington’s proprioceptive field being referred to as proprioceptors, the term has also been used for the mechanoreceptors located at the surface of the body, and portions of the vestibular apparatus responsible for conveying information regarding the orientation of the head with respect to gravity. Thus, to avoid potential confusion from this wide disparity of use, we recommend utilizing more specific references to the mechanoreceptors of interest.

Neuromuscular control is a frequently used term in many disciplines related to motor control. It can refer to any of the aspects surrounding nervous system control over muscle activation and the factors contributing to task performance. Specifically, from a joint stability perspective, we define neuromuscular control as the unconscious activation of dynamic restraints occurring in preparation for and in response to joint motion and loading for the purpose of maintaining and restoring functional joint stability. Although neuromuscular control underlies all motor activities in some form, it is not easily separated from the neural commands controlling the overall motor program. For example, in throwing a ball, particular muscle activation sequences occur in the rotator cuff muscles to ensure that the optimal glenohumeral alignment and compression required for joint stability are provided. These muscle activations take place unconsciously and synonymously with the voluntary muscle activations directly associated with the particulars of the task (ie, aiming, speed, distance). Proprioceptive information concerning the status of the joint and associated structures is essential for neuromuscular control. The use of proprioception for motor control and neuromuscular control is the focus of part II of this article.

PERIPHERAL SENSORY PATHWAYS

Sources of Proprioceptive Input

Based on Sherrington’s definition of the proprioceptive field, the mechanoreceptors responsible for proprioceptive information are primarily found in muscle, tendon, ligament, and capsule with the mechanoreceptors located in the deep skin and fascial layers traditionally associated with tactile sensations being theorized supplementary sources. In general, mechanoreceptors are specialized sensory receptors responsible for quantitatively transducing the mechanical events occurring in their host tissues into neural signals.
necessarily occurs in a similar manner across the various mechanoreceptors, each morphologic type possesses some degree of specificity for the sensory modality to which it responds (light touch versus tissue lengthening), as well as the range of stimuli within a sensory modality. \(^{31}\) As several detailed reviews have been published on the subject, \(^{11,22-24,28,32-34}\) we offer only a brief review of the characteristics and functions of joint and muscle mechanoreceptors.

Although 4 types of receptors are dispersed throughout ligamentous and capsular tissues, Ruffini receptors are the most frequently described. \(^{22}\) They are considered to behave as both static and dynamic receptors based on their low-threshold, slow-adapting characteristics. \(^{26}\) In contrast, the low-threshold, rapidly adapting characteristics of Pacinian corpuscles cause them to be exclusively classified as dynamic receptors. \(^{26}\) Also present in these tissues are Golgi tendon organ-like endings and free nerve endings. \(^{11,26,28,35}\)

Mechanoreceptors located within musculotendinous tissue include the Golgi tendon organs (GTOs) spaced along the musculotendinous junction at varying intervals and the muscle spindles located in the muscle tissue. Through each GTO passes a small bundle of muscle tendon fibers destined to attach to muscle fibers. This series arrangement, coupled with the very low threshold and high dynamic sensitivity exhibited by the sensory endings, enables GTOs to provide the CNS with feedback concerning muscle tension. \(^{23}\) GTOs function primarily in signaling active muscle tension (tension developed during contraction) rather than passive tension (tension developed during inactive muscle stretching). \(^{23}\)

As a whole, muscle spindles are responsible for conveying information regarding muscle length and rate of changes in length. Muscle spindles consist of specialized afferent nerve endings that are wrapped around modified muscle fibers (intrafusal fibers), several of which are enclosed in a connective tissue capsule. \(^{19,36}\) There are different types of intrafusal fibers: some are mainly sensitive to changes in muscle length, whereas others are more sensitive to the rate of change in muscle length. \(^{36}\)

Although the central areas of the intrafusal muscle fibers lack contractile elements, the peripheral areas contain contractile elements, which are innervated independent of extrafusal (skeletal) muscle fibers via the gamma motor neurons (γ MNs). Activation of the peripheral contractile elements stretches the central regions containing the sensory receptors from both ends. This results in an increase in the firing rates of the sensory ending and an increase in the sensitivity of the muscle spindle to length changes. \(^{19}\) At the spinal level, various peripheral receptors, such as skin receptors, articular receptors, and chemoreceptors, strongly influence the activity of the γ-MN system \(^{24,37-41}\) and, therefore, the muscle spindle in providing afferent information.

**Sensory Integration at the Spinal Cord Level**

Integration of sensory input received from all parts of the body is largely considered to begin at the level of the spinal cord. Integration describes the summation, gating, and modulation mechanisms that occur as a result of various combinations of excitatory and inhibitory synapses with the afferent neurons. \(^{7}\) These synapses may originate from several sources, such as other afferent fibers or neurons conveying descending signals from higher CNS structures. Afferent integration is an essential component of coordinated, fluid motor control and occurs along all levels of the CNS. This section offers only a brief overview of afferent integration at the spinal level, as a detailed review has previously been published. \(^{42}\)

In contrast to the few tactile neurons that travel directly to the cortex without synapsing, \(^{43}\) many of the axons conveying proprioceptive information bifurcate once they enter the dorsal horn of the spinal cord to synapse with interneurons. The essence of afferent integration at the spinal level lies with the interneurons and the neurons connecting with higher CNS levels. Control over these neurons via descending commands from the brainstem and cortex provides these centers with the ability to filter the sensory input that will be conveyed via the ascending tracts. \(^{7}\) In other words, the supraspinal CNS regions modulate the sensory information from the periphery that enters the ascending tracts.

An additional hypothesis, the final common input hypothesis proposed by Johansson et al. \(^{24}\) presents an additional and supplemental integrative mechanism. This hypothesis resides on the strong influence that the muscle, skin, and joint afferents and descending pathways have over gamma neuron activation. \(^{24}\) As mentioned previously, the peripheral regions of intrafusal muscle fibers contain contractile elements innervated by γ MNs, with the level of activation directly controlling muscle spindle sensitivity. Any of the signals barraging the γ-MN pools alter their level of activation, and, therefore, influence the input arising from the muscle spindles. Thus, the afferent signals from muscle spindles are hypothesized to be a function of muscle length changes superimposed on the integrated peripheral receptor and descending pathway information. In this manner, the γ-MN system may be considered a “premotor neuronal integrative system” that conducts “polymodal feedback” to the CNS (Figure 3). \(^{24}\)

**Proprioceptive Coding to Higher CNS Centers**

Two theories describe the methods by which specific proprioceptive messages from the various receptors are conveyed to the CNS. The first theory, the labeled line theory, is based
on the presumption that each unique stimulus triggers a certain receptor connected to a specific nerve fiber that terminates at a specific point or multiple points within the CNS. Critics of this theory suggest that it neglects the fact that most receptors and neurons appear to be sensitive to different types of stimuli and not only to a specific stimulus. The second theory, ensemble coding, suggests that proprioceptive information is transferred to the CNS through an encoding across a neural population of receptors rather than discrete units from the individual receptors. Originally proposed by Erickson, this theory proposes that receptors possess unique, but overlapping, ranges of sensitivity. Application of this theory to the sensorimotor system has been largely a result of the work by Johansson et al. Clinically, this theory may help explain the improved conscious proprioceptive acuity and reduction in subjective instability complaints associated with elastic wraps and neoprene bracing.

**Ascending Spinal Tracts Conveying Proprioceptive Information**

Most proprioceptive information travels to higher CNS levels through either the dorsal lateral tracts or the spinocerebellar tracts. The 2 dorsal lateral tracts are located in the posterior region of the spinal cord and ultimately convey the signals to the somatosensory cortex. Although the majority of the sensations traveling in this tract are touch, pressure, and vibration, various amounts of the conscious appreciation of position and kinesthetic sensations have also been attributed to this tract. The spinocerebellar tracts are characterized by the fastest transmission velocities in the CNS. As their name suggests, the spinocerebellar tracts terminate in various areas of the cerebellum, where the signals may be processed and integrated with other afferent and descending information. In contrast to the conscious sensory appreciation associated with the dorsal lateral tracts, the spinocerebellar tracts are believed to be responsible for “nonconscious proprioception” (ie, limb position, joint angles, and muscle tension and length) used for reflexive, automatic, and voluntary activities. In addition to relaying peripheral afferent information, parts of these tracts are associated with transmitting an effenter copy of motor neuron drive back to the higher CNS levels.

**Conscious Perception of Proprioception**

Sherrington’s early 1900s view attributing the sense of kinesthesia and joint position sense (“muscular sense”) to muscle receptors was accepted for most of the century, with a brief hiatus existing for a short time period (1950–1970) when several authors considered joint receptors to be the primary source. The change of belief was initiated by the results of several studies considering occulomotor system problems and the overall lack of evidence supporting direct group I afferent projections to the sensorimotor cortex. The premise shifted back to muscle receptors after the demonstration of joint receptors’ response voids through the midranges of motion and reports of movement illusions caused by tendon vibration. Our survey of the available literature on this topic up to present times reveals a plethora of conflicting evidence supporting each tissue’s receptors (joint, muscle, and cutaneous) as the predominant source. Even more uncertain is supposition on the contribution individual morphologic receptors make within each tissue (joint, muscle, and cutaneous) during functional, full-range joint movements. Rather than attempting to review all the original work conducted in this area, which by itself would become a lengthy paper, we will highlight some of the major findings and discuss the implications of the continued controversy with respect to conscious appreciation of joint position sense (JPS) and kinesthesia.

The first step in determining whether a group of tissue receptors could potentially contribute to conscious appreciation of kinesthesia and JPS is through documentation of response sensitivity through the full physiologic range of joint motion. Through the use of animal models, several investigators have concluded that mechanoreceptors located in the joint capsule do not appear to be sufficiently stimulated through the midranges of motion to contribute substantially to proprioception, especially in relation to the seemingly potent input stemming from muscle receptors. Several authors have concluded, based on this evidence, that joint capsular afferents are unlikely to signal JPS and kinesthesia information through the midranges of motion and that their only proprioceptive function is signaling endranges of motion. Grigg discredited ligamentous receptors as probable candidates based on their low numbers (with respect to joint capsule) and inability to signal specific joint movement and position. It is important to note, however, that the evidence upon which many of these conclusions are based was collected during passive movements. As Pedersen et al. stated, researchers have reported increases in joint receptor working ranges (angular range in which a receptor remains active) during active movements. Similar to joint afferents, cutaneous afferents have been speculated to respond only at the extremes of motion. Unfortunately, this finding is not without controversy, as several authors have recently attributed cutaneous mechanoreceptors with a precise ability to convey joint movements through skin strain patterns. In contrast to joint and cutaneous mechanoreceptors, muscle spindles have been almost universally described as able to respond unidirectionally across the entire physiologic range of movement.

As mentioned previously, proprioception for conscious appreciation travels via the dorsal lateral tracts, with the contributions to these tracts from muscle and joint mechanoreceptors remaining largely unknown. Thus, demonstrating the existence of projections to the cortical sensory areas and conscious perceptions after direct receptor stimulation is the second necessity in determining the predominant source of conscious proprioception (Figure 4). Unfortunately, the results of these studies have complicated the conclusions one would draw based solely on the sensitivity evidence. Cortical projections have been reported from joint (both capsular and ligament) afferents, muscle spindles, and GTOs. With respect to conscious appreciation of peripheral receptor stimulation, electric stimulation of both joint and cutaneous (slowly adapting type II) afferent fibers were reported to elicit sensations related to the relevant joint and evoke perceptions of joint movement, respectively. Edin and Johansson demonstrated that mechanical stimulation of cutaneous receptors elicited kinesthetic sensations. While direct stimulation of a single muscle spindle afferent failed to elicit movement perception, stimulation of several muscle spindles through vibration and isolated traction has been reported to evoke conscious movement sensations. The failure of joint and cutaneous afferents anesthesia to disrupt conscious kinesthesia and JPS provides further support for the importance of muscle receptors in conscious proprioception.
LEVELS OF MOTOR CONTROL

The motor components of the sensorimotor system contributing dynamic joint stability are synonymous with areas controlling whole-body motor control. These components consist of a central axis and 2 associate areas. The central axis corresponds to the 3 levels of motor control, spinal cord, brain stem, and cerebral cortex, whereas the 2 associate areas, cerebellum and basal ganglia, are responsible for modulating and regulating the motor commands. Sensory information underlies the planning of all motor output and, as described in previous sections, is conveyed to all 3 levels of motor control. Activation of motor neurons may occur in direct response to peripheral sensory input (reflexes) or from descending commands initiated in the brain stem or cerebral cortex, or both. Independent of the initiating source, skeletal muscle activation occurs through signal convergence onto the motor neurons located in the spinal ventral horns. This concept is what Sherrington labeled the final common path. Both types of motor neurons, alpha motor neurons (α MNs) controlling extrafusal muscle fibers (skeletal) and γ MNs controlling intrafusal muscle fibers (muscle spindles), exit the spinal ventral horns.

The central axis areas are organized in both a hierarchic and parallel manner. The hierarchic organization allows the lower motor areas to automatically control the details of common motor activities, while the higher centers can devote resources to controlling the more precise and dexterous motor activities. In addition, as mentioned earlier, higher levels can regulate the afferent information reaching them through inhibitory and facilitatory control over sensory relay nuclei. Through the parallel arrangement, each motor control center can directly issue independent contributory descending motor commands directly on the motor neurons.

Spinal Cord Level

It should be apparent from our earlier discussion that the spinal cord plays an integral role in motor control, despite the gross anatomy suggesting it may only be a medley of conduction pathways. From the spinal cord arise direct motor responses to peripheral sensory information (reflexes) and elementary patterns of motor coordination (rhythmic and central pattern generators). As discussed earlier, very little afferent input and few descending commands synapse directly on motor neurons. Instead, most input terminates upon the interneurons located throughout all areas of cord gray matter. Even in the case of a simple monosynaptic reflex, such as the stretch reflex, bifurcations from the incoming afferent fiber arise. These bifurcations may convey the afferent information to a number of locations, including interneurons, higher motor centers, and other motor neurons (antagonistic). The bifurcations and interneuronal networks provide the basis for the spinal cord’s efferent integrative functions.

Reflexes may be elicited from the stimulation of cutaneous, muscle, and joint mechanoreceptors and may involve excitation of α MNs, γ MNs, or both. For many clinicians, the stretch reflex in response to rapid muscle lengthening provides the most familiar example. These reflexes, as well as the other reflexes attributed to the spinal cord neuronal circuitry, are more complex than simply direct input-output connections. Superimposed on even the simplest monosynaptic reflexes are influences from such sources as other afferent input, descending commands, or both.

Brain Stem

Despite being the most primitive part of the brain from a phylogenetic perspective, the brain stem contains major circuits that control postural equilibrium and many of the automatic and stereotyped movements of the body. In addition to being under direct cortical command and providing an indirect relay station from the cortex to the spinal cord, areas of the brain stem directly regulate and modulate motor activities based on the integration of sensory information from visual, vestibular, and somatosensory sources.

Two main descending pathways, the medial and lateral pathways, extend from the brain stem to the spinal cord neural networks. The medial pathways influence the motor neu-
rons innervating the axial and proximal muscles, while the lateral pathway controls the distal muscles of the extremities. In addition to controlling postural control, some axons comprising the medial pathways make excitatory and inhibitory (including suppression of spinal reflexes) synapses with the interneurons and motor neurons involved with movement and postural control. Through influences on the γ MNs, parts of both the medial and lateral tracts assist in maintaining and modulating muscle tone.

Cerebral Cortex

In general, the motor cortex is responsible for initiating and controlling more complex and discrete voluntary movements. It is divided into 3 specialized and somatotopically organized areas, each of which project directly and indirectly (via the brain stem) onto interneurons and motor neurons located in the spinal cord.74 The first area, the primary motor cortex, receives peripheral afferent information via several pathways and is responsible for encoding the muscles to be activated, the force the recruited muscles produce, and the direction of the movement.43,72 The second area, the premotor area, also receives considerable sensory input72; however, it is mainly involved with the organization and preparation of motor commands. The supplemental motor area, the third specialized area of the motor cortex, also plays an important role in programming complex sequences of movement that involve groups of muscles.72,74

The major direct descending pathway from the motor cortex to the α MNs and γ MNs is the corticospinal tract. In addition to influencing motor functions directly, the corticospinal tract also affects motor activity indirectly through the descending brain stem pathways.

Associate Areas

Although the 2 associate areas, the cerebellum and basal ganglia, cannot independently initiate motor activity, they are essential for the execution of coordinated motor control. The cerebellum, operating entirely at a subconscious level, plays a major role in both the planning and modification of motor activities though comparison of the intended movement with the outcome movement.75,76 This is accomplished through the continuous inflow of information from the motor control areas and the central and peripheral sensory areas. The cerebellum is divided into 3 functional divisions. The first division receives vestibular input, both directly and indirectly from the vestibular labyrinth (semicircular and otolith receptors) and, as might be surmised based on the input, is involved with postural equilibrium. The second cerebellar division is mainly responsible for the planning and initiation of movements, especially those requiring precise and rapid dexterous limb movements.75 This division receives input from both the sensory and motor cortices. It is the third division, the spinocerebellum, which receives the somatosensory information conveyed through the 4 ascending spinocerebellar tracts. In addition to the somatosensory input, this division of the cerebellum also receives input from the vestibular labyrinth and visual and auditory organs. The output from the spinocerebellum serves to adjust ongoing movements through influential connections on the medial and lateral descending tracts in the brain stem and cortex via projections on the vestibular nucleus, reticular formation, red nucleus, and motor cortex.75 In addition to controlling movements, the spinocerebellum also uses the somatosensory input for feedback regulation of muscle tone through regulation of static γ-MN drive to the muscle spindles.75 Lastly, the cerebellum also receives an efferent copy of the motor commands arriving at the ventral roots of the spinal cord.76 The cerebellum has also been implicated in motor learning.7,75

The basal ganglia consist of 5 subcortical nuclei (groups of nerve cells) located deep within the cerebral hemispheres. In contrast to the cerebellum, which has input and output connections with all 3 levels of motor control, the cerebral cortex is the only central axis component having input and output connections (via the thalamus) with the basal ganglia.43,77 With respect to motor control, the basal ganglia are believed to be involved with more higher-order, cognitive aspects of motor control.77 An additional distinction from the cerebellum is that the basal ganglia receive input from the entire cerebral cortex, not just those associated with sensory and motor function.77 The widespread input and output cortical connections suggest that they are involved with many functions other than motor control.

CONCLUSIONS

The sensorimotor system encompasses all of the sensory, motor, and central integration and processing components involved with maintaining joint homeostasis during bodily movements (functional joint stability). We have attempted to introduce the physiology of joint stability through an in-depth presentation of the sensorimotor system. As evident from the sections concerning ascending proprioception pathways and levels of motor control, the sensorimotor system is much more complex than a simple input-output system that resides primarily in the lower levels of motor control. Rather, activation of the dynamic restraints, and therefore, functional joint stability, arises from components synonymous with the entire motor control system of the body. Thus, functional joint stability is an inherently complex and complicated physiologic process. In the absence of mechanical stability, the fact that many individuals return to preinjury levels suggests that some degree of compensatory mechanisms can be developed to provide the supplemental stability required. These compensatory mechanisms most likely arise from the dynamic restraints of the involved joint, as well as motor adaptations at proximal and distal segments. This would suggest the importance of the supraspinal temporal and spatial organization of the dynamic restraint activation. In part II of this article, we will discuss the importance of proprioception in organizing muscle activation for both motor control and sensorimotor control of functional joint stability.

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