Opinion
Phantom Limbs, Neuroprosthetics, and the Developmental Origins of Embodiment
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Amputees who wish to rid themselves of a phantom limb must weaken the neural representation of the absent limb. Conversely, amputees who wish to replace a lost limb must assimilate a neuroprosthetic with the existing neural representation. Whether we wish to remove a phantom limb or assimilate a synthetic one, we will benefit from knowing more about the developmental process that enables embodiment. A potentially critical contributor to that process is the spontaneous activity – in the form of limb twitches – that occurs exclusively and abundantly during active (REM) sleep, a particularly prominent state in early development. The sensorimotor circuits activated by twitching limbs, and the developmental context in which activation occurs, could provide a roadmap for creating neuroprosthetics that feel as if they are part of the body.

Our Bodies, Our Selves
Embryonic limbs begin as small buds protruding from the body wall. Over time, these buds differentiate into limbs that enable walking, running, reaching, grasping, hugging, and touching. To support such diverse functions, fully developed limbs are highly complex systems composed of multiple joints set in motion by distinct muscles, a proprioceptive sense provided by muscle spindles and Golgi tendon organs, and diverse receptors for touch and vibration [1]. Moreover, the machinery of the limb is integrated at many levels with the even more complex neural circuitry of the spinal cord and brain. When this process of integration is done well, we accept each limb as a part of us. We sense that we ‘own’ it. This sense of body ownership and the neural machinery that supports it constitutes a body schema (see Glossary) [2].

Inherent in the notion of a body schema is the intertwining of mind and body. In contrast with the view that the mind can be understood without consideration of the body, an embodied perspective emphasizes the critical contributions to mind of such factors as sensory–motor coupling, body morphology, and physical constraint [3]. In other words, although minds control aspects of the body, bodies also shape the mind [4,5].

Today, the concept of embodiment is informing the creation of biologically inspired robots [6]. But even the most sophisticated robots still do not experience one of the most foundational of all biological processes – development. Although development may not seem like a particularly important process for a robot, the central idea behind the emerging field of developmental robotics is that a contemporary robot – precisely because it does not develop – can never achieve the flexibility and adaptability required for life in a growing, ever-changing body [5,7–9].

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Accordingly, for a robot to be as flexible and adaptable as a human, it must experience a developmental process.

If development can inform the creation of better robots, then perhaps it can also inform our understanding of other similar problems, including efforts aimed at integrating synthetic and biological body parts. But for development to be a guide for understanding these sorts of problems, we must first understand the process by which developing limbs become incorporated into the body schema.

Sleep, Myoclonic Twitching, and Embodiment

Long before limbs are used for moving infants around a room or reaching for objects, they move spontaneously and with a seeming lack of purpose. From the earliest stages of embryonic development, spontaneous movements are necessary for normal limb development, including bone, cartilage, and joint formation. Such movements can be caused by factors within the muscle (i.e., myogenic) or from signals provided by the spinal cord and brain (i.e., neurogenic); over development, myogenic movements fade as neurogenic movements gain in importance.

Two types of neurogenic movement emerge over the course of development. The first type occurs when infants are awake and comprises large, continuous, or sustained movements of the limbs, such as stretching, kicking, and yawning. The second type of movement occurs during active sleep, which is most prevalent early in development, and comprises discrete, brief, and jerky movements of the limbs, digits, face, eyes, and, in rodents and many other mammals, whiskers and tail. These movements are called myoclonic twitches. Twitching continues into adulthood in select muscle groups, depending on species-typical morphology and functional importance.

Traditionally, twitching was considered a by-product of a dreaming brain or a sign of incomplete motor inhibition. The former view is negated by an absence of evidence for cortical involvement – and an abundance of evidence for brainstem involvement – in the production of twitches. The latter view has more merit, but cannot easily explain the persistence of twitching into adulthood in various muscle groups across a diverse array of species. In contrast, when considered in its own right within the context of development, we find that twitching limbs exhibit complex spatiotemporal patterns across multiple joints, reflecting the emergence of motor synergies. But for twitching to actually contribute to – rather than merely reflect – the development of the sensorimotor system, it is necessary that the twitch movements themselves trigger sensory feedback and that this feedback, in turn, activates sensorimotor structures.

Sensorimotor processing depends on a complex network of structures at each level of the neuraxis, and now substantial evidence supports the idea that reafference from twitching limbs triggers neural activity at each of those levels: from the spinal cord to the dorsal column nuclei (which include the external cuneate nucleus), cerebellar cortex and deep cerebellar nuclei, thalamus, cerebral cortex, and hippocampus. (Activity in the basal ganglia is also likely to be influenced by twitch-related reafference, but this possibility has not yet been investigated.) Moreover, the red nucleus, which is part of a group of premotor structures in the mesencephalic junction, contributes to the production of twitches while also receiving reafference, suggesting that the integration of motor output and sensory feedback occurs even within this so-called motor nucleus. It also notable that the nuclei within the mesencephalic junction are associated with the evolution of limbs and flapping fins, and appear to have been differentially recruited across species for the control of highly specialized structures such as the human hand and the...
Figure 1. Twitches Are Produced Discretely and Are Spatiotemporally Organized. (A) Sequential high-speed video frames from an 8-day-old rat to show the discrete nature of twitching, illustrated here using the left elbow (top) and right shoulder (bottom). The yellow arrows indicate the direction in which the limb is moving and the white markers were used for motion tracking of limb movements. (B) Records of limb twitching in an 8-day-old rat at a long (top) and short (bottom) timescale to illustrate the spatiotemporal organization of twitching; the segment denoted by the gray box is expanded below. Each tick mark indicates a single twitch at the shoulder, elbow, or wrist in the right (red) or left (blue) forelimb. Unbroken and dashed lines indicate the direction of movement at each joint (i.e., flexion or extension, adduction or abduction). Adapted from [16].

Box 1. Myoclonic Twitching across Species and across the Lifespan

Because twitching has traditionally been interpreted as a by-product of other brain processes (see [63]), there has been little motivation to assess the quantity and patterning of twitching in infants or adults [16]. But without basic descriptive and quantitative data about twitching, we cannot fully evaluate its possible significance.

To help remedy this situation, we created a website to serve as a clearing-house for video recordings of sleep-related twitching in all its forms. The website links to the multitudes of videos available on YouTube from people across the globe who are fascinated by the behavior of sleeping animals (including humans) at home, in zoos, and in the wild. The website also includes videos donated by sleep scientists and laypeople. Although just beginning, the site contains numerous videos representing 40 different species, from aardvarks to wild boars, and even some insects, across the lifespan.

What emerges from viewing this collection of videos is the sheer diversity and ubiquity of twitching. Also, certain patterns begin to emerge. For example, although twitching is similarly abundant in the limbs and face of kittens, puppies, piglets, ferret kits, and rat pups, twitching is more heterogeneous in adults, with some body parts twitching much more than others. This seems particularly clear in those species that have highly specialized morphologies. For example, twitches of the eyes (i.e., rapid eye movements) occur in humans, dogs, capybaras, and many other species, but so do twitches of the whiskers in cats and rats, the snout in pigs, the digits of ferrets, the ears of bats, the claws of aardvarks, the neck of giraffes, and the trunk of elephants. Thus, in addition to the general maintenance of sensorimotor function in adults, twitching may be especially important for the calibration of those specialized parts of the body that are involved in active sensing [64]—that is, the process by which animals direct movable sensors toward objects of interest so as to maximize the precision of stimulus detection, enhance perception, and achieve behavioral ends with greater efficiency.
elephant’s trunk [33]. Twitching, it seems, is enmeshed within the neural machinery that gives limbs their purpose.

Just as the red nucleus is more than merely a motor nucleus, the functions of motor cortex extend beyond motor control. We have seen this quite clearly by recording neural activity in this region of cortex in sleeping infant rats and noting the prominent neural activation that follows the onset of even the most delicate twitch. In marked contrast, during wake movements in the same pups – and even as the limbs move vigorously – the motor cortex is often silent [28]. These observations led to two conclusions. First, the motor cortex plays a negligible role in the production of movement, whether sleep or wake movement, at these early ages. This conclusion is wholly consistent with what is known about the brainstem generators of twitches [32] and the development of the cortical control of wake movements [34].

Second, the motor cortex begins its existence primarily as a recipient and processor of sensory input. In fact, even in adults, the sensory function of motor cortex is well established [35] (see also [36] for evidence that sensory cortex plays a role in motor control). In infants, the sensory drive to motor cortex is more salient because its motor function has yet to emerge. Moreover, the relative absence of reafferent activation of motor cortex during wake movements suggests

Figure 2. Circuit Diagram Illustrating the Web of Neural Structures That Support Embodiment. (A) Some major sensorimotor structures involved in the production of movement and the processing of reafferent signals. Reafference from twitching limbs triggers neural activity within all of the sensorimotor loops shown: these loops include afferent and efferent projections from nuclei in the MDJ (red), cerebellum (blue), and sensorimotor cortex (green). Importantly, the limb receives motor commands from both motor cortex and nuclei in the MDJ (including the red nucleus), and the reafferent pathways complete each loop, allowing each structure to monitor and update its representation of the limb. (B) A phantom limb is perceived when the limb is absent, but the sensorimotor structures representing the limb remain intact. Without reafference to update the limb’s neural representation, the phantom limb persists. (C) A closed-loop neuroprosthetic limb interfaces directly with motor cortex and sensory cortex (yellow arrows), providing reafference that bypasses subcortical sensorimotor structures. By interacting exclusively or primarily with sensorimotor cortex, the rest of the existing limb representation is bypassed, thereby limiting the incorporation of the neuroprosthetic limb into the body schema. Abbreviations: IO, inferior olive; LRN, lateral reticular nucleus; MDJ, mesodiencephalic junction.
The presence of a state-dependent gating mechanism that passes twitch-related reafference but blocks wake-movement-related reafference. We recently identified such a gating mechanism in a medullary structure, the external cuneate nucleus, which receives direct proprioceptive input from muscles in the forelimbs and neck [20]. The demonstration of a mechanism that preferentially passes twitch-related reafference throughout the infant brain supports the idea that twitching represents a unique mechanism for developing and refining sensorimotor circuits and somatotopic maps (see Box 2).

One critical upshot of the foregoing discussion is that the motor functions of the brain, including motor cortex, are built on a foundation of sensation. Thus, sensorimotor integration is a concept that goes to the heart of what behavioral competencies require: a complex network of inextricable links within and among sensorimotor – not sensory and motor – structures. Thus, whatever a body schema is, it is the product of the highly distributed network illustrated in Figure 2A, including the sensorimotor cortex, cerebellum, and associated structures. But also, the body schema is the product of a developmental process through which limbs are fully assimilated with the nervous system. By this account, twitches both enable the development of embodiment and provide a window onto the architecture of embodiment.

**Mimicking Development to Enhance Recovery of Sensorimotor Function**

Because neural plasticity is so clearly heightened in early development, one key to rewiring the adult brain is to ‘remove the brakes’ on plasticity [37]. But development is characterized not only by the absence of brakes on plasticity, but also the presence of spontaneous activity – in the visual, auditory, and sensorimotor systems – that promotes it [38]. If twitching contributes to developmental plasticity in the sensorimotor system, then perhaps we can use this knowledge to heighten plasticity in adults.
As one striking example of developmental plasticity, consider the corticorubral tract, which connects the motor cortex to the ipsilateral red nucleus. In response to unilateral damage to the motor cortex in infant rats, the undamaged motor cortex not only retains its connections with the ipsilateral red nucleus, but also sprouts new connections with the contralateral red nucleus, thus giving the undamaged cortex control over both sides of the body. Critically, this spontaneous rewiring occurs only if the cortical damage occurs before 10 days of age in rats. Given that this period of heightened plasticity corresponds with the developmental period of maximal active sleep and twitching, it may be that twitching directly contributes to both the typical development of the corticorubral system and its ability to rewire after early injury.

Although rewiring of the adult rat corticorubral system does not occur spontaneously after injury, recent studies nonetheless show that rewiring is possible. For these studies, the brain damage entails unilateral transection of the corticospinal tract at the level of the medulla, which causes profound unilateral deficits in forelimb control. To promote recovery after the transection, the investigators electrically stimulated the unaffected motor cortex for 6 hours a day for 10 days. This protocol induced sprouting of connections from the stimulated motor cortex to the contralateral red nucleus and other structures, once again allowing one side of the brain to gain control over both sides of the body. Moreover, forelimb motor function was restored.

For the present purposes, what is most striking about this protocol is how the researchers carefully calibrated their stimulation protocol so as to induce discrete movements of the elbow, wrist, or digits of the forelimb; notably, stimulations occurred during the day when rats spend most of their time asleep. Accordingly, it is possible – although still untested – that this stimulation protocol, by inadvertently mimicking some aspects of twitching, owes its effectiveness to its ability to discretely activate both motor and sensory circuits involved in forelimb control.

It should be noted that a stimulation protocol that more closely mimics twitching would entail stimulation of the red nucleus and other brainstem premotor nuclei that are known to produce twitching; however, these brainstem nuclei are much less accessible than motor cortex. But such details may not matter so long as the stimulation protocol results in the activation of sensorimotor circuits throughout the neuraxis in a discrete and somatotopically precise way.

### Phantom Limb as a Window onto the Origins of Embodiment

The sensorimotor system’s heightened developmental plasticity is powerfully illustrated by the behavioral feats of humans born without limbs – from hand-walking in people born without legs to the remarkably dexterous feet and toes of so-called ‘armless wonders’. These individuals exemplify how humans and other animals learn to use the body they have, not the body they were ‘supposed’ to have. Although the developmental plasticity inherent in the sensorimotor system enables functionality in individuals with atypically formed bodies, it is also essential for the everyday process of adapting to the increases in limb size, shape, and strength that characterize typical development across a diversity of species, including those with extreme morphological and behavioral specializations.

When a limb is removed, either through accident or surgery, there remains in over 90% of amputees a vivid sense that the limb remains exactly where it was. Such phantom limbs are fascinating, but they are also instructive about the neural circuits that are devoted to each limb and make it possible for the limb to function as part of an integrated whole. As illustrated in Figure 2B, the loss of a limb leaves behind these neural circuits, thereby providing the material basis for the phantom. Thus, to accomplish a truly comprehensive amputation, one would have...
to remove not only the limb, but also the web of limb-devoted sensorimotor structures that, in their entirety, encompass the schema for that limb.

Phantom limb most commonly arises in adults after trauma or surgery. Moreover, the likelihood of phantom limb depends on the age at which a limb was lost. According to one study [44], limb amputation at 4–6 years of age produced a phantom in 91% of cases, whereas the incidence dropped to about 62% at 2–4 years of age and to 33% at 0–2 years of age; moreover, the incidence was still 20% in those children born with missing limbs (see also [45,46]).

The fact that phantom limb arises in cases of congenital limb loss – no matter how low the incidence – led some to conclude that the neural system that enables phantom limb is, at least in part, innate or, in other parlances, hardwired, genetically determined, or preprogrammed [44,47–49]. However, as is typically the case with such claims of innateness [50], empirical evidence argues for an alternative, developmental explanation.

Ten years ago, in a reexamination of phantom limb associated with early limb loss, Price [51] proposed that limbs are incorporated into the body schema through a process that begins in utero with ‘spontaneous muscle activity.’ He suggested that spontaneous limb activity gives rise to sensory feedback, thereby driving the development of the neural networks that underlie the perception of a phantom limb. Although this proposal did not consider a role for sleep and sleep-related twitches (indeed, the proposal focused on spontaneous movements that are ‘initiated within the muscles themselves’), it is easy to imagine how twitches could lay the foundation for a phantom limb in the minority of cases involving congenital amputation – especially given the fact that the active sleep is even more abundant in preterm than full-term human infants [52]. Accordingly, the odds of developing a phantom may depend on whether a fetus’s limb failed to develop at all or was amputated (e.g., due to amniotic band syndrome [53]) after having developed and begun to move.

Beyond Repair: Fulfilling the Promise of Neuroprosthetics

The ultimate test of our understanding of embodiment is playing out today within the burgeoning field of brain–machine interfaces (BMI). Fulfilling BMI’s promise of achieving the complete assimilation of a neuroprosthetic limb with an existing nervous system will require advances in many fields, including computational neuroscience, sensor technology, and materials science [54]. It is also possible that this promise will depend upon the incorporation of knowledge about the developmental process by which natural limbs are assimilated.

Researchers widely recognize that only a closed-loop BMI, in which the motor control of a neuroprosthetic limb is accompanied by some form of sensory feedback, will achieve complete assimilation [54,55]. To close the loop, sensory feedback from vision can be used to adjust the movements of a prosthetic toward a goal. But to achieve maximum functionality and a stronger sense of assimilation, diverse sensory feedback – especially proprioceptive and tactile feedback – will likely be critical [56,57].

As illustrated in Figure 2C, closed-loop neuroprosthetic control is most readily achieved using implanted devices that translate signals from motor cortex into output commands to a prosthetic and convey sensory signals from the prosthetic to sensory cortex [54,55]. However, this approach fails to engage most of the sensorimotor network normally involved in limb control.

Therefore, as prosthetic sensor technology improves, the incorporation of a neuroprosthetic limb into the body schema may require maximal engagement of the cortical and subcortical components that constitute the sensorimotor system. To accomplish this, sensory input from
the neuroprosthetic might be routed into the nervous system through peripheral nerves (e.g., in the stump) or through the spinal cord or brainstem. Moreover, drawing inspiration from development and the stimulation experiments described earlier, a brain stimulation protocol that drives movement in the neuroprosthetic and triggers associated sensory feedback throughout the sensorimotor system may be most effective for inducing plasticity and enabling somatotopic remapping.

Concluding Remarks
Developmental scientists emphasize the daily quantity of input that enables and supports the development of such skills as walking [58] and word learning [59]. Twisting fits squarely within this perspective: the many thousands of discrete twitches that are produced each day by the human infant provide ample opportunity for the activity-dependent development of the sensorimotor system and the assimilation of growing limbs into the nervous system.

The ideas put forward here about the possible value of leveraging understanding of development, sleep, and twitching to improve recovery of function after injury and improving neuroprosthetics are speculative. The ideas rest on the simple fact that the muscles and sensory receptors of developing limbs establish functional connectivity at every level of the sensorimotor system from the spinal cord to the forebrain. In addition, sleep-related twitches provide ample opportunity for limbs to form those functional connections via discrete activations during active sleep, a state that provides an ideal low-noise environment for the transmission of high-fidelity signals. Whether these considerations can help to improve the assimilation of neuroprosthetics into the nervous system of children or adults remains an open question (see Outstanding Questions).

Oliver Sacks observed that all amputees understand “that a phantom limb is essential if an artificial limb is to be used” (p. 66) [60]. In other words, the neural machinery that produces a phantom is the same machinery with which a fully functional neuroprosthetic must be assimilated. Our claim here is that limbs assimilate with the nervous system through a developmental process, and that a fuller understanding of that process will help us design more lifelike neuroprosthetics. In turn, improvements in neuroprosthetic technology will afford new opportunities for addressing basic questions in sensorimotor control [61], including questions about the roles that active and quiet (non-REM) sleep play in the consolidation of motor memories [62]. Ultimately, this interplay between basic science and advanced technology will help us answer some of the most ancient questions concerning the origins of our sense of self and the mechanisms that sustain it.

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Resources
www.twitchsleep.net

References

Outstanding Questions
Does reafference from twitches contribute to the plasticity that occurs after tasks or manipulations that engage the sensorimotor system, such as limb immobilization, finger-sequence learning, and the wearing of eye prisms?

Is it possible to enhance recovery of sensorimotor function in humans after stroke or injury through the use of therapies that explicitly mimic twitching? If so, which features of twitching (e.g., discrete stimulation, occurrence during active sleep, site of stimulation) are most effective for fostering recovery?

Which structures or circuits within the sensorimotor system are most important for modifying a body schema? Studies of phantom limb and neuroprosthetic assimilation in brain-damaged populations may help to answer this question.

Can sleep – and sleeping with a neuroprosthetic limb – enhance the assimilation of that limb into an existing nervous system? How does brain activity during active and quiet (non-REM) sleep change throughout the process of assimilation?

Do individual differences in the ability to assimilate a neuroprosthetic relate to individual differences in sleep and twitching?

The likelihood that a phantom limb arises after an amputation increases through the first several postnatal years. Do individual differences in sleep and twitching help to explain individual differences in the occurrence of a phantom limb?