We are IntechOpen, the world's leading publisher of Open Access books Built by scientists, for scientists

6,900

186,000

200M

Download

154
Countries delivered to

Our authors are among the

TOP 1%

most cited scientists

12.2%

Contributors from top 500 universities



WEB OF SCIENCE

Selection of our books indexed in the Book Citation Index in Web of Science™ Core Collection (BKCI)

Interested in publishing with us? Contact book.department@intechopen.com

Numbers displayed above are based on latest data collected.

For more information visit www.intechopen.com



The Integrative Role of the Basal Ganglia

Clemens C.C. Bauer, Erick H. Pasaye, Juan I. Romero-Romo and Fernando A. Barrios

Additional information is available at the end of the chapter

http://dx.doi.org/10.5772/54189

1. Introduction

Emergent experimental and clinical evidence supports the notion that the cortico-basal ganglia-thalamo-cortical loops progress along parallel circuits connecting cortical and subcortical regions subserving the processing of sensorimotor information, associative and affective knowledge [1]. In particular the role of the basal ganglia has long been known to be involved in motor control because of the marked deficits associated with their damage. However, the exact aspects of motor control that they have under normal conditions have not been clear at all. The traditional view is that the basal ganglia are involved in the selection and inhibition of action commands [2], but an increasing number of brain-imaging studies show that the basal ganglia, besides being involved in motor tasks are also involved in more integrative and cognitive processes such as mental imagery [3,4], sensory processing [5,6], planning [7], attention [8,9], and language [6,10,11]. This evidence supports the view that the basal ganglia output not only targets the primary sensory-motor cortices, but also specific areas of premotor and prefrontal cortex, which include the oculomotor area of the cortex, the dorsolateral prefrontal cortex, lateral orbitofrontal cortex, and anterior cingulate/medial orbitofrontal cortices [12]. Thus, having the ability to influence not only sensory-motor control, but also several different types of cognitive and limbic affective functions [12] which underlie complex and integrative processes such as self-awareness, introspective perspective of one's own self and consciousness [13]. This integrative role between lower afferent input and higher integrative and executive stages of information processing require an intact and closed loop of information flow to generate the primary experience of self and thus self-agency. Self-agency has tentatively been defined as the feeling of being the author of one's own actions [14]. Thus, when we move our arm, we know (a) that it is our arm and (b) that it is us, who moves the arm. One approach to understand the complex integration of afferent and efferent information processing and the integration in the self is the internal model theory of motor control [15]. According to this theory there are two functionally different components in the motor system, inverse and



forward circuits. It is assumed that inverse models provide the motor commands necessary to achieve a desired consequence of an action, specified by higher-level goals (e.g., intentions). One is fully aware of the desired consequences of an action, but unaware of the motor programs per se. Forward models predict the sensory consequences of each motor program to be executed, an idea known as the efference copy, a model first put forward in von Holst and Mittelstaedt [16] and which has been extended in recent years with the "null" hypothesis of Ramachandran [17]. Accordingly, it is claimed that, whenever a motor program is issued, an efference copy is produced in parallel, this is, a prediction of the sensory consequences expected after the execution of the program based on exactly this efference copy. The internal model theory of motor control has been successfully applied to explain a whole variety of disorders related to the awareness of actions [for a review see 17] but the role played by the basal ganglia in this model has not been very clear. Here we follow this model extending it to the performance of a complex cognitive task, such as mental imaginary movement of a limb and the coexistent conscious awareness of just imagining it and actually refraining from moving it which involves exactly this subtle combination between the forward model requiring intact peripheral efferent/afferent information pathways and the inverse model requiring intact higher-level cortical areas which include the basal ganglia-thalamus-cortex pathway [19]. Moreover, we contrast the normal integrity of this forward/inverse model loop in healthy subjects with an abnormal open loop in amputees where an essential part of the loop has been disrupted. We thus argue that because of this abnormal and open loop involving the basal ganglia and the thalamocortical system the conscious awareness of the phantom phenomenon is created.

2. Problem statement

Waking up from anesthesia, an amputee faces the conflict in the experience of self, between the conscious vividness of his phantom limb (PL) and the lack of correlation with reality [20–23]. In the urge to discover whether he was actually amputated, the patient looks under the sheet for visual self-recognition and is, in a flash, confronted with this new reality of an absent limb. The resulting cognitive conflict between the seen embodiment and the felt one, in most cases carries on resulting in the perception of a ghost of their amputated limb as a phantom [24]. Giummarra et al. [20] report phantom limb experiences that include phenomena of (a) perception of bodily aspects of phantom limbs such as size (in relation to the intact limb), shape, posture, and telescoping (or shortening) of the phantom; (b) exteroceptive and proprioceptive sensations and (c) prosthesis embodiment. Early studies of phantom limb movement were carried out using combined techniques of EEG, MEG, and fMRI in order to locate its representation in the sensory-motor cortex [25-28] and in the cerebellum [29,30]. Later studies described the distinct functional anatomy of the mental representation of imaginary movements [31], during planning, visualizing, and motor intention [32], both in healthy subjects as well as in patients with different neural diseases. Other studies have centered on the difference between imaginary movement and executed movement [3,33]. A recent study by Diers et al. [34] showed activation in the supplementary motor area (SMA) cortex after PL imagined movement. And in a more recent study Pasaye et al. [35] have described nuerocorrelates of the PL perception using fMRI. Since the first conceptualization of the phenomenon of phantom limb in amputees in 1915, as the manifestation of the persistence of the body schema of the missing limb [36], many theories have arisen. Most of these theories suggest a central role of reorganization of the somatosensory cortex, others advocate for a perceptual completion [37], or the product of conflicting cues from central reorganizational changes [38]. Although some degree of sensory or motor impairment is constant, recent models of motor awareness suggest that the reduction in afferent information may be less critical than higher-level reorganization related to the subjective correlates of action planning and motor intention [15,39]. In contras, a number of studies of patients with amputated limbs encourage that a conscious perception of a body part by tactile stimulation does not necessarily require the integrity or even existence of the tactile receptors on the skin, or the body parts themselves. That is, the neural representation of the body in the brain is sufficient to elicit an awareness of the body part or tactile stimuli in the absence of its physical counterpart (e.g., the limbs themselves) [40]. We believe that the reduction or absence in afferent information is a key factor that in combination with higher-level reorganization generates the conscious awareness of the phantom phenomenon. Here we hypothesize that for the performance of a complex cognitive task, such as imaginary movement of a limb, and the concomitant conscious awareness of just imagining it and actually refraining from moving it, an involvement of a subtle combination between intact peripheric (efferent/afferent) information pathways and intact higher-level cortical areas (basal-ganglia-thalamus-cortex pathway) are needed [19]. In amputees, this normal combination of lower- and higher-level processes is disrupted and could be the underlying cause of conscious awareness of the phantom limb. This arrangement of processes was reported by Staub et al. [41], who found an increased blood oxygen level dependent (BOLD) signal in the Basal Ganglia-Thalamic-Motor-Cortex loop pathway during imaginary movement of a patient with chronic supernumerary phantom limb, which developed only in association with motor intent directed at a hemiplegicanesthetic upper limb. Staub's finding is analogical to what Ramachandran proposes as the "null" signal hypothesis in the mirror neuron system (MNS), which prevents activity in the MNS from reaching the threshold for conscious awareness [17] or related to what Fitzgibbon et al. [42] suggests as the underlying cause in synesthesia for pain. We think that the logic behind Ramachandran's hypothesis is plausible, and if applied to an intact afferent somatosensoryproprioceptive and efferent motor feedback, no activity in the basal-ganglia-thalamus-cortex loop would be seen in healthy subjects, in contrast to the incomplete closing of the afferent/efferent loop and therefore an absence of the "null" signal in the case of amputees. Hence the inhibition of activity in the basal ganglia-thalamus-cortex pathway is disrupted, thereby generating an abnormal open loop functioning of the thalamocortical system and its consequent activation. This abnormal activation of the thalamocortical system, we suggest, could be the underlying cause of the conscious awareness of the phantom phenomenon.

This hypothesis is also in accordance with an emergent change of view in the functionality of the basal ganglia, from the classic view of being just part of the common motor pathway, to a more integrative, dynamic and resource-selection mechanism that participates in the sensorymotor, affective, and cognitive process related to the executive planning and selection of an action mechanism [43]. Furthermore, the conscious awareness and the subsequent sense of agency [44] which patients report to have had as they perform imagined tasks with their amputated limbs has never between contrasted with healthy subjects.

In the current study, we tested the hypothesis that the basal ganglia-thalamus-cortex pathway is disrupted in amputees, as compared to control subjects, and that this disruption is the key to the cascade of conscious awareness of the phantom limb. We contrasted between lower limb amputees and control subjects as they performed a simulated neurocognitive motor-imagery task with their phantom toes or intact toes respectively.

3. Method used

3.1. Subjects

Six unilateral lower limb amputees, 3 with left lower limb (LLL) and 3 with right lower limb (RLL) amputation (mean age 35.3, range 15-60 years, for details see Table 1), and 6 healthy controls (HC, mean age 29.16, range 20-59 years) participated in the study. All participants gave written informed consent prior to taking part in the study and the local institutional review board approved the protocol, which adhered to the Declaration of Helsinki. None of the subjects had neurological or psychiatric disorders.

	Age ¹ Ctrl	Age ² Pat	Amp ³ side	Amp ⁴ site	Cause of ⁵ Amputation	SP ⁶	PLP ⁷	PLS ⁸
1	20	34&	RLL	TF	Traumatic	+	-	+
2	21	15	RLL	TT	Traumatic	+	-	+
3	22	44	RLL	TT	Traumatic	+	-	+
4	24	60	LLL	TT	Traumatic	+	-	+
5	29	33	LLL	TF	Traumatic	+	-	+
6	59	26	LLL	TT	Traumatic	+	-	+

Table 1. Clinical data description of controls and patients. All subjects are right handed males. ¹Column for control subjects' age. ²Column for patients' age. ³Three patients had right lower limb amputation and three had left lower limb amputation.⁴Site of surgery. ⁵Car or train accident was main cause of amputation. ⁴All the patients reported stump pain (SP), but 7none ever had Phantom Limb Pain (PLP). ⁸All patients were able to move at will their phantom toes (extension/flexion).

3.2. Experimental procedure

The experiment consisted of two parts. In the 'executed movement' condition the participants were instructed to make a flexion/extension movement with the intact (amputees) or non-dominant (HC) toes. In the condition 'imagined movements' the amputees imagined making a flexion/extension movement with the phantom toes and the HC imagined the same movement with their dominant toes. For all conditions the subjects had their eyes open. All movements or

imagined movements were observed under close scrutiny by the researcher. All conditions were separate blocks of fMRI measurements with durations of 30 seconds each, separated by resting periods of 30 seconds. Each condition was repeated three times. There was no training session for either the control subjects or the amputees.

3.3. fMRI measurement

The fMRI scans were conducted with a GE 1.5 T GE LX Magnetic Resonance instrument (Milwaukee, WI, USA) using the standard quadrature headcoil. Subject's head was securely fastened in the head holder to minimize movement. Functional images were acquired with GE EPI-BOLD pulse sequence with 90° flip angle TE=60 ms, TR=3000 ms, over seven contiguous coronal sections, 8.0 mm thick with zero gap. Imaging was centered near the central gyrus. Structural images localized exactly over the same seven sections of the functional studies were obtained using a high resolution T1 weighted protocol. The activation was done in boxcar block paradigm of 30 seconds of stimulated state and 30 seconds of un-stimulated state, over a total of three blocks each.

3.4. Image analysis

All MR data was transferred to an offline workstation all images were translated into timeordered stacks using the software MRIcro (Chris Rorden, http://www.cabiatl.com/mricro/). The experiments were analyzed subject by subject following a standard motion correction with image registration, to do a more precise alignment, since the study was acquired in coronal slices, the high resolution T1-weighted images were aligned to the T1-weighted SPM mask in MNI space, the transformations were saved and applied to the functional EPI time ordered stacks, and then the functional images were normalized to a ROI of the EPI standard provided by SPM, spatial smoothing using a Gaussian kernel of FWHM 6mm and high pass temporal filtering. Functional signal was obtained with a block model convolved with an hemodynamic response function (HRF) without time derivative correction these single subject analysis resulted in contrast maps used in the second level analysis. The second level analysis was executed to estimate group average activation using a Student-t maps limited and adjusted with p = 0.05 with no Volterra interactions.

4. Results

Analysis of the data was carried out for each subject individually, to see if there were statistically significant activation clusters, and then by group after combining all subjects. The average functional maps obtained revealed ipsilateral or contralateral brain activation sites, which were colored according to the tasks performed on each lower limb of all subjects: green-colored brain activation sites correspond to the right leg, while red-colored brain activation sites correspond to the left leg. Yellow-colored sites correspond to areas of overlapping activation during the performance of the tasks on each limb. The results of the three groups' brain activation sites are summarized in Table 2.

		Tala	irach c	oordinates	RIGHT		Talairach coordinates		
BRAIN REGION	BA	x	y z		BRAIN REGION	BA	x	y	Z
MIN T	-00	<i>(</i> 1	27 (Left amputee, executing right imaginary motion	Will T	0.1	F1	20	_
Middle Temporal Gyrus	22	-61	-37 6		Middle Temporal Gyrus	21	51	-29	-5
Paracentral Lobule	4	-10	-38 63						
Medial Frontal Gyrus	6	-6	-14 62						
Precentral Gyrus	6	-30	-20 64	Į.					
Lentiform Nucleus,	*	-26	-8 4						
Putamen				Left amputee, executing					
Medial Frontal Gyrus	6	-8	-11 50	left virtual motion	Medial Frontal Gyrus	6	2	-26	64
Sub-Gyral	37	-48	-39 -5		Medial Frontal Gyrus	6	6	-9	61
Superior Temporal Gyrus	22	-59	-35 9		Substania Nigra	*	8	-24	-14
Thalamus	*	-8	-25 1		Superior Temporal Gyrus	21	48	-27	-5
Precentral Gyrus	4	-42	-11 47	7					
				Right amputee, executing left imaginary motion					
Superior Temporal Gyrus	22	-65	-42 15	i	Superior Temporal Gyrus	22	50	-4	-1
Superior Temporal Gyrus	22	-65	-18 -1		Superior Temporal Gyrus	22	50	-12	1
Postcentral Gyrus	3	-44	-17 54		Medial Frontal Gyrus	6	8	-11	50
Cingulate Gyrus	24	-8	-10 41		Middle Frontal Gyrus	6	40	-5	46
					Cingulate Gyrus	24	8	-12	37
					Thalamus	*	18	-9	13
				Right amputee, executing right virtual motion					
Precentral Gyrus	4	-12	-32 62		Superior Temporal Gyrus	22	51	-4	-3
Medial Frontal Gyrus	6	-10	-26 58	3					

		Talairach coo			ordinates	RIGHT		Talairach coordinates		
BRAIN REGION	BA	x	y	z		BRAIN REGION	BA	x	y	Z
Superior Temporal		-63								
Gyrus										
Superior Temporal	*	-63	-21	3						
Gyrus										
	4	-50	-12	41						
Lentiform Nucleus,		-16	-10	-6						
Medial Globus										
Pallidus										
Substania Nigra	*	-8	-10	-10						
					Control,					
					executing left imaginary					
					motion					
Inferior Frontal	45	-61	20	16		Medial Frontal	6	8	-12	71
Gyrus						Gyrus				
						Medial Frontal	6	2	-3	61
-						Gyrus				
					Control,					
					executing right					
					imaginary motion					
Medial Frontal	6	-2	3	51		Postcentral Gyrus	43	65	-16	21
Gyrus										
1	6	-8	-6	68						
Gyrus										
Medial Frontal	6	-8	1	53						
Gyrus				_						
Superior Temporal	22	-50	4	2						
Gyrus BA=Brodmann Area										

BA=Brodmann Area.

Talairach coordinates: x (left[-], right[+]); y (posterior[-], anterior[+]), z (inferior[-], superior[+]). * no Brodmann area related

Table 2. Anatomical location of activation clusters during imaginary and virtual motion.

The average functional maps obtained from LLL amputee during the imaginary movement of the toes of both feet also present distinct cortical and subcortical activity. Performance of imaginary movement of the toes of the right intact toes showed activation sites bilaterally at the STG (BA 21,22), contralateral interhemispheric M1 (BA 4), contralateral SMA (BA 6), and contralateral Putamen. During the performance of imaginary movement with the left amputated toes, distinct cortical and subcortical activities were observed at the following sites: bilateral interhemispheric SMA (BA 6), bilateral STG (BA 21,22), ipsilateral M1 (BA 4), ipsilateral Subgyral (BA 37), ipsilateral Thalamus, and contralateral Substantia Nigra (figure 1,A).

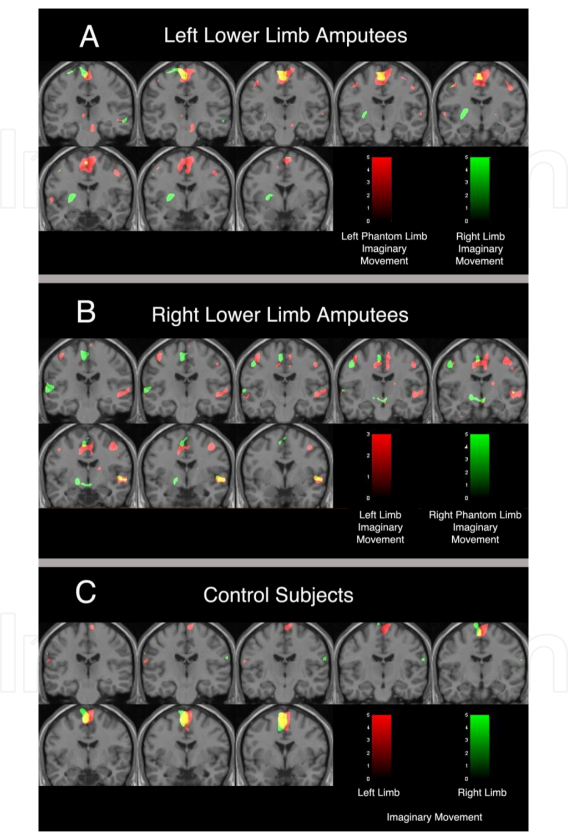


Figure 1. Average functional activation maps during imaginary movement of the right leg (in green) and left leg (in red) for A) left lower limb amputees, B) right lower limb amputees and C) control subjects. All images are presented in radiological convention.

Average functional activation maps acquired from RLL amputees during the imaginary movement of the toes of both feet depict both cortical and subcortical activities. During the performance of imaginary movement of the toes of the left intact toes, there is bilateral Temporal activity (BA 22), and bilateral Anterior Cingulate Cortex (ACC, BA 24), ipsilateral Primary Somatosensory Cortex (SI, BA 3), contralateral SMA (BA 6), and contralateral Thalamic activity, while during the performance of imaginary movement with the right amputed toes, there is a distinct activation, namely of the bilateral STG (BA 42, 22), contralateral Primary Motor Cortex (M1, BA 4), contralateral SMA (BA 6), contralateral Basal Ganglia at the Medial Globus Pallidus and at the Substantia Nigra (figure 1, B).

The average functional maps obtained from the six control subjects as they performed imaginary movement with (a) the toes of their right leg show activated contralateral sites corresponding to interhemispheric Supplementary Motor Area (SMA, BA 6), Superior Temporal Gyrus (STG, BA 22), and Ipsilateral Postcentral Gyrus (IPG, BA 43); while the performance of the same tasks with (b) the toes of their left leg, activated similar contralateral interhemispheric SMA (BA 6) and ipsilateral prefrontal (PF, BA 45) brain areas (figure 1,C).

5. Discussion

Here we compared the brain activations of imagined and executed movements of the intact toes and phantom toes in lower limb amputees, with the imagined and executed movement of the toes of healthy controls, using fMRI. Both, patients and control subject expressed that they initially had to exert greater effort in this self-generating dual process of 1) trying to resolve an apparent conflict between the simultaneous intent to move their toes and refraining from moving them or closing the sensory-motor feedback loop; and 2) locating, by means of imagery monitoring, a distant portion of the body-image, which is an attention/memory task. This sensation can be do to the increased contribution of the prefrontal cortex, in particular the dorsolateral prefrontal cortex (DLPFC), which is known to participate in motor imagery, not just in the sensory-motor integration and the attention/memory neurocognitive task with the anterior cingulate cortex, but also in a joint route with the posterior parietal cortex (PPC) during motor imagery [30,45-47]. The PPC, as a multisensory integrative cortex, plays an important role in the cognitive dynamics for spatial representation (limb-position) and movement intent, attention, working memory, and guidance of action [48,49]. As Jeannerod proposed "If motor imagery occurred with execution deliberately blocked or delayed, the representation would be protected from cancellation and would become accessible to conscious processing" [50]. Additionally, controls and amputees reported that they were consciously aware of their intact toes or, in the case of amputees, that they had the conscious perception of their phantom toes during the imaginary task. The brain activations show similarities but also differences between amputee groups during the imaginary movement tasks of their intact and amputated toes. The differences were: First, both amputee groups activate Basal Ganglia areas during the performance of the imaginary movement of the amputated toes. Second, the RLL amputee group shows more lateralized brain activation than the LLL amputee group. Third, during

the imaginary movement task, the RLL amputee group seems to require a greater attention control (ACC) as they performed the imaginary movement with their left intact toes than the LLL amputee group with their right intact toes. Additionally, the brain activations observed in the amputee group during the imagery movement task of the amputated toes involved the Basal Ganglia loop (RLL amputee group = Lentiform Nucleus, Medial Globus Pallidus, and Susbtantia Nigra; LLL amputee group = Thalamus and Substantia nigra). Thus, the imagined movement task in amputees demands different circuitry subsets for its accomplishment, namely: 1) the attention/memory/guidance loop, 2) the kinesthetic imagery loop, and 3) the conflict intention loop. The computational logistics for such activity can only be carried out by means of the intracortical and cortical-subcortical loops between Thalamus and Basal Ganglia nuclei. The kinesthetic representation of an action or a planned motor intent is the combined result of a widely distributed neuronal ensemble between DLPFC, inferior frontal cortex, and the SMA [31], together with the posterior parietal cortex (PPC) and the ACC for spatial awareness, attention and multisensory integration [49,51]. It is of mayor importance to notice here that the activation of the Basal Ganglia loop was not seen during imagery movement task of the intact toes in amputees or the healthy control group. As far as the motor intent is concerned (Fig 1, B): a) in the control group, the performance of the task with the left (non-dominant) toes activated contralateral SMA (BA 6) and ipsilateral inferior frontal cortex (BA 45), while the same task with the right (dominant) toes activated contralateral SMA (BA 6), STG (BA 22), and contralateral postcentral gyrus (BA 43). The minimal brain activity found in the controls' kinesthetic representation correlates with the ensemble proposed [52,53]. b). In the RLL amputee group, in the coronal volumes (Fig 1, A), the performance of the imaginary motor intent with the left intact toes activated the bilateral ACC (BA 24), STG (BA 22), ipsilateral S1 (BA 3), and broad contralateral interhemispheric activity from SMA (BA 6) and contralateral thalamus. However, during the right imagined movement task of the amputated toes, there is bilateral activation at STG (BA 22/42), contralateral SMA (BA 6), M1 (BA 4), contralateral Lentiform Nucleus (medial globus pallidus), and Substantia Nigra. The RLL amputees' brain activity for the kinesthetic representation differed between the performance of the intact imaginary movement and the amputated imaginary movement, since Lentiform Basal Ganglia activity is only present during the amputated toes imaginary movement. c) In the coronal-volumes of the LLL amputee group (Fig 1, C) during the performance of the imaginary movement task with the right intact toes, besides a bilateral MTG (BA 21/22) activation, there is clear contralateral Lentiform Nucleus-Putamen activation, together with SMA (BA 6) and M1 (BA 4). However, during the imagery movement task with the amputated toes of the left leg, there is an ipsilateral thalamic and a Subgyral (BA 37) activation and a contralateral Substantia Nigra activation besides the large bilateral SMA (BA 6) and a bilateral MTG (BA 21/22) activations observed during the intact toes imaginary movement. The LLL amputee brain activity for the kinesthetic representation differed slightly between the performance of the imaginary movement of intact versus amputated toes, since there is Basal Ganglia activity (Putamen) during imaginary movement of the intact toes, while during amputated toes imagined movement there is Thalamic and Substantia Nigra activity. This Basal Ganglia-Thalamo-Motor-Cortex loop subserves several cortex functions, such as memory tasks, orientation in space, and the ability to change behavioral set [43,54]. In the motor

imagery task, in particular in LL amputee research literature, the role of this gangliathalamo-motor-cortex loop has never been mentioned. In this study we set out to establish its presence using the already mentioned task in the amputee and control groups. Thus, by comparing control and amputee groups, we found that there is minimal cortical activation difference between them, however, the difference occurs in the subcortical activation of the Basal Ganglia loop, since in both, the control group and during intact toes imaginary movement of amputated subjects there is no Basal Ganglia activation, while the activation of distinct Lenticular-Substantia-Nigra-Basal-Ganglia-Thalamic loop is clear in the amputee group performing imaginary movement of the amputated toes. We thus propose that the recruitment of these Basal Ganglia plays an important role in the process of conscious awareness of a missing limb reported by amputees.

It is important to point out that we set out to find the involvement of the Basal Ganglia-Thalamic-Motor-Cortex loop by means of this motor imagery task, as part of our hypothesis that the amputee can and does move the phantom limbs at will, this, do to his framework of body awareness as part of a self-related neurocognitive experience that can be as diverse as the perceiving of size, shape, posture, itch, touch, pressure, vibration, temperature, 'electric' sensations and prosthesis embodiment and has been well documented [20]. Similar to Ramachandran's "null" hypothesis [17] we think that the interruption of the thalamic afferences/efferences may explain the persistence of an open loop functioning of the thalamocortical system and its consequent activation, which is a key factor to the cascade of conscious awareness and stability of the phantom phenomenon. This open loop functioning of the thalamocortical system is revealed in the present study by the increased blood oxygen level-dependent (BOLD) signal in the Basal Ganglia-Thalamic-Motor-Cortex loop pathway during imaginary movement of the amputated toes. Thus, supporting our hypothesis of the abnormal closed-loop functioning of the thalamocortical system as underlying the phantom phenomenon.

6. Conclusion

To conclude, we have put forward evidence of amputee patients' indirect responses to PL experiences for an objective evaluation that suggests that the conscious awareness of a phantom limb emerges from both the reduction in afferent information and the higher-level brain reorganization of the cognitive representations of the amputee's own body. We based our assumptions on the hypothesis that the thalamocortical loop is closed in healthy subjects, which enable them to distinguish an imaginary movement as actually being just imagined. This, do to the feedback received from intact peripheric (efferent/afferent) information pathways. The evidence shown here thus suggests that this abnormal open loop of the basal-ganglia-thalamocortical system underlies the conscious awareness of the phantom limb. The current approach further suggests that the basal ganglia within this basal-ganglia-thalamocortical system loop play a crucial and complex integration of afferent and efferent information processing. Furthermore, this integration creates the conscious awareness of the self and is in line with the internal model theory of motor control [15] where inverse and forward models of information processing interact continuously and reciprocally. The inverse model component in the motor system providing the motor commands necessary to achieve a desired consequence of an action, specified by higher-level goals and the forward model predicting the sensory consequences of each of these motor programs to be executed. Accordingly, whenever a motor program is issued, an efference copy is produced in parallel and an accurate prediction of the sensory consequences expected after the execution of the program, which in turn informs the inverse program of the actual state of the self and closing the loop for the next command. With this normally closed loop, the integration of the self is achieved and a normal body ownership and awareness is crated which is necessary to create the autobiographical experience of self.

Glossary of terms

Bottom-up: direction of information flow from the periphery (i.e. sensory cells or mechanoreceptors) to the central nervous system.

Top-down: information flow from central nervous system toward peripheral effector cells (i.e. muscles).

Efferent: Conveying away from the central nervous system

Afferent: Conveying towards a central nervous system

Phantom limb: is the sensation that an amputated or missing limb is still attached to the body

Somatosensory system: sensory system composed of the receptors and processing centers to produce the sensory modalities such as touch, temperature, proprioception (body position), and nociception (pain).

Proprioception: sensory modality that processes the body position

Ipsilateral: same side of the body **Contralateral:** other side of the body

Author details

Clemens C.C. Bauer*, Erick H. Pasaye and Fernando A. Barrios Neurobiology Institute, Universidad Autónoma de México, Querétaro, México

Juan I. Romero-Romo

Querétaro General Hospital, SESEQ; Queretaro, Mexico

Acknowledgement

We want to thank Dr. Perla Salgado and Dr. Rafael Rojas form the Imaging Department of The ABC Medical Center, Mexico City. This work was partially supported by CONACyT R31162-A, and a Doctoral Schollarship from CONACyT México and the Doctoral Program in Biomedical Sciences in the Institute of Neurobiology, Universidad Nacional Autónoma de

^{*} Corresponding Author

México for CCCB and EHP. We thank Leopoldo Gonzalez-Santos, Juan J. Ortiz for their technical support.

7. References

- [1] Draganski B, Kherif F, Klöppel S, Cook PA, Alexander DC, Parker GJM, et al. (2008) Evidence for Segregated and Integrative Connectivity Patterns in the Human Basal Ganglia. Journal of Neuroscience. 9;28(28):7143-52.
- [2] Alexander GE, DeLong MR, Strick PL. (1986) Parallel organization of functionally segregated circuits linking basal ganglia and cortex. Annual Review of Neuroscience. 9(1):357-81.
- [3] Lotze M, Montoya P, Erb M, Hülsmann E, Flor H, Klose U, et al. (1999) Activation of cortical and cerebellar motor areas during executed and imagined hand movements: an fMRI study. Journal of Cognitive Neuroscience. 11(5):491–501.
- [4] Lorey B, Bischoff M, Pilgramm S, Stark R, Munzert J, Zentgraf K. (2009) The embodied nature of motor imagery: the influence of posture and perspective. Experimental Brain research. 194(2):233-43.
- [5] Mizumori SJ., Puryear CB, Martig AK. (2009) Basal ganglia contributions to adaptive navigation. Behavioural brain Research. 199(1):32-42.
- [6] Kotz SA, Schwartze M, Schmidt-Kassow M. (2009) Non-motor basal ganglia functions: a review and proposal for a model of sensory predictability in auditory language perception. Cortex. 45(8):982-90.
- [7] Bernsz S, Sejnowskil TJ. (2010) How the basal ganglia make decisions. Available: http://papers.cnl.salk.edu/PDFs/How%20the%20Basal%20Ganglia%20Make%20Decisio ns%201996-2876.pdf. Accesed 2012 March
- [8] van Schouwenburg MR, den Ouden HE., Cools R. (2010) The human basal ganglia modulate frontal-posterior connectivity during attention shifting. The Journal of Neuroscience. 30(29):9910-8.
- [9] Provost JS, Petrides M, Monchi O. (2010) Dissociating the role of the caudate nucleus and dorsolateral prefrontal cortex in the monitoring of events within human working memory. European Journal of Neuroscience. 32(5):873-80.
- [10] Friederici AD, Kotz SA. (2003) The brain basis of syntactic processes: functional imaging and lesion studies. Neuroimage. 20:S8-S17.
- [11] Booth JR, Wood L, Lu D, Houk JC, Bitan T. (2007) The role of the basal ganglia and cerebellum in language processing. Brain Research. 1133:136-44.
- [12] Middleton FA, Strick PL. (2000) Basal ganglia and cerebellar loops: motor and cognitive circuits. Brain Research Reviews. 31(2-3):236-50.
- [13] Kircher TTJ, Leube DT. (2003) Self-consciousness, self-agency, and schizophrenia. Consciousness and Cognition. 12(4):656–69.
- [14] Lycan WG. (1996) Consciousness and Experience. A Bradford Book.
- [15] Frith CD, Wolpert DM. (2000) Abnormalities in the awareness and control of action. Philosophical Transactions of the Royal Society of London. Series B: Biological Sciences. 355(1404):1771-88.

- [16] Holst E, (1950) Mittelstaedt H. Das reafferenzprinzip. Naturwissenschaften. 37(20):464–
- [17] Ramachandran VS, Brang D. (2009) Sensations evoked in patients with amputation from watching an individual whose corresponding intact limb is being touched. Archives of Neurology. 66(10):1281.
- [18] Kircher T, David AS. (2003) The self in neuroscience and psychiatry. Cambridge Univ Pr.
- [19] Serino A, Haggard P. (2010) Touch and the body. Neuroscience & Biobehavioral Reviews. 34(2):224-36.
- [20] Giummarra MJ, Georgiou-Karistianis N, Nicholls ME., Gibson SJ, Chou M, Bradshaw JL. (2010) Corporeal awareness and proprioceptive sense of the phantom. British Journal of Psychology. 101(4):791–808.
- [21] Hunter JP, Katz J, Davis KD.(2008) Stability of phantom limb phenomena after upper limb amputation: a longitudinal study. Neuroscience. 156(4):939–49.
- [22] Halligan PW. (2002) Phantom limbs: the body in mind. Cognitive Neuropsychiatry. 7(3):251-69.
- [23] Ramachandran VS, Hirstein W. (1998) The perception of phantom limbs. The DO Hebb lecture. Brain. 121(9):1603.
- [24] Arzy S, Thut G, Mohr C, Michel CM, Blanke O. (2006) Neural basis of embodiment: distinct contributions of temporoparietal junction and extrastriate body area. The Journal of Neuroscience. 26(31):8074.
- [25] Schreiber A, Ball T, Kristeva-Feige R, Mergner T, Feige B, Scheremet R, et al. (1998) Primary and Secondary Motor Areas in fMRI and EEG. Available: http://cds. ismrm.org/ismrm-1998/PDF6/p1568.pdf. Accessed 2012 March
- [26] Stippich C, Freitag P, Kassubek J, Sörös P, Kamada K, Kober H, et al. (1998) Motor, somatosensory and auditory cortex localization by fMRI and MEG. Neuroreport. 9(9):1953.
- [27] Flor H, Elbert T, Mühlnickel W, Pantev C, Wienbruch C, Taub E. (1998) Cortical reorganization and phantom phenomena in congenital and traumatic upper-extremity amputees. Experimental Brain Research. 119(2):205-12.
- [28] Condés-Lara M, Barrios FA, Romo JR, Rojas R, Salgado P, Sánchez-Cortazar J. (2000) Brain somatic representation of phantom and intact limb: a fMRI study case report. European Journal of Pain. 4(3):239–45.
- [29] Nitschke MF, Kleinschmidt A, Wessel K, Frahm J. (1996) Somatotopic motor representation in the human anterior cerebellum. A high-resolution functional MRI study. Brain: Brain (1996) 119 (3): 1023-1029. doi: 10.1093/brain/119.3.1023.
- [30] Ersland L, Rosén G, Lundervold A, Smievoll AI, Tillung T, Sundberg H. (1996) Phantom limb imaginary fingertapping causes primary motor cortex activation: an fMRI study. Neuroreport. 8(1):207.
- [31] Jeannerod M, Frak V. (1999) Mental imaging of motor activity in humans. Current Opinion in Neurobiology. 9(6):735–9.
- [32] Ganis G, Thompson WL, Kosslyn SM. (2004) Brain areas underlying visual mental imagery and visual perception: an fMRI study. Cognitive Brain Research. 20(2):226-41.

- [33] Lotze M, Flor H, Grodd W, Larbig W, Birbaumer N. (2001) Phantom movements and pain An fMRI study in upper limb amputees. Brain. 124(11):2268.
- [34] Diers M, Christmann C, Koeppe C, Ruf M, Flor H. (2010) Mirrored, imagined and executed movements differentially activate sensorimotor cortex in amputees with and without phantom limb pain. Pain. 149(2):296-304.
- [35] Pasaye E, Gutiérrez R, Alcauter S, Mercadillo R, Aguilar-Castañeda E, de Iturbe M, et al. (2010) Event-Related Functional Magnetic Resonance Images During the Perception of Phantom Limb; a Brushing Task. The Neuroradiology Journal. 23:665–70.
- [36] Pick A. Zur Pathologie des Bewußtseins vom eigenen Körper. (1915) Ein Beitrag aus der Kriegsmedizin. Neurologisches Zentralblatt. 34:257-65.
- [37] Levine DN. (1990) Unawareness of visual and sensorimotor defects: A hypothesis. Brain and Cognition. 13(2):233-81.
- [38] Ramachandran VS, Hirstein W. (1998) The perception of phantom limbs. The DO Hebb lecture. Brain. 121(9):1603.
- [39] Heilman KM. (1991) Anosognosia: possible neuropsychological mechanisms. In Prigatano, G. P., & Schacter, D. L. (1991). Awareness of deficit after brain injury: Clinical and theoretical issues. pp 53-62. Oxford University Press, USA.
- [40] Gallese V. (2007) The "conscious" dorsal stream: embodied simulation and its role in space and action conscious awareness. Psyche. 13(1):1–20.
- [41] Staub F, Bogousslavsky J, Maeder P, Maeder-Ingvar M, Fornari E, Ghika J, et al. (2006) Intentional motor phantom limb syndrome. Neurology. 67(12):2140-6.
- [42] Fitzgibbon BM, Giummarra MJ, Georgiou-Karistianis N, Enticott PG, Bradshaw JL. (2010) Shared pain: from empathy to synaesthesia. Neuroscience & Biobehavioral Reviews. 34(4):500-12.
- [43] Gurney K, Prescott TJ, Wickens JR, Redgrave P. (2004) Computational models of the basal ganglia: from robots to membranes. Trends in Neurosciences. 27(8):453-9.
- [44] Ramachandran VS. (1993) Behavioral and magnetoencephalographic correlates of plasticity in the adult human brain. Proceedings of the National Academy of Sciences of the United States of America. 90(22):10413.
- [45] Decety J, Perani D, Jeannerod M, Bettinardi V, Tadary B, Woods R, et al. (1994) Mapping motor representations with positron emission tomography. Nature. 371, 600-602
- [46] Stephan KM, Fink GR, Passingham RE, Silbersweig D, Ceballos-Baumann AO, Frith CD, et al. (1995) Functional anatomy of the mental representation of upper extremity movements in healthy subjects. Journal of Neurophysiology. 73(1):373–86.
- [47] Boussaoud D. (2001) Attention versus intention in the primate premotor cortex. Neuroimage. 14(1):S40-S45.
- [48] Culham JC, Kanwisher NG. (2001) Neuroimaging of cognitive functions in human parietal cortex. Current Opinion in Neurobiology. 11(2):157-63.
- [49] Andersen RA, Buneo CA. (2002) Intentional maps in posterior parietal cortex. Annual Review of Neuroscience. 25(1):189–220.
- [50] Jeannerod M. (1994) The representing brain: Neural correlates of motor intention and imagery. Behavioral and Brain Sciences. 17(02):187–202.

- [51] Crammond DJ. (1997) Motor imagery: never in your wildest dream. Trends in Neurosciences. 20(2):54.
- [52] Jackson PL, Decety J. (2004) Motor cognition: a new paradigm to study self-other interactions. Current Opinion in Neurobiology. 14(2):259-63.
- [53] Jeannerod M, Decety J. (1995) Mental motor imagery: a window into the representational stages of action. Current Opinion in Neurobiology. 5(6):727–32.
- [54] Yang TT, Gallen CC, Ramachandran VS, Cobb S. (1994) Noninvasive detection of cerebral plasticity in adult human somatosensory cortex. Neuroreport: An International Journal for the Rapid Communication of Research in Neuroscience. Vol 5(6), 701-704.

