Apraxia. A Review.

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Abstract

Praxic functions are frequently altered following brain lesion, giving rise to apraxia, a complex pattern of impairments that is difficult to assess or interpret. In this chapter, we review the current taxonomies of apraxia and related cognitive and neuropsychological models. We also address the questions of the neuroanatomical correlates of apraxia, the relation between apraxia and aphasia and the analysis of apraxic errors. We provide a possible explanation for the difficulties encountered in investigating apraxia and also several approaches to overcome them, such as systematic investigation and modeling studies. Finally, we argue for a multidisciplinary approach. For example, apraxia should be studied in consideration with and could contribute to other fields such as normal motor control, neuroimaging and neurophysiology.

Introduction

Apraxia is generally defined as “a disorder of skilled movement not caused by weakness, akinesia, deafferentation, abnormal tone or posture, movement disorders such as tremor or chorea, intellectual deterioration, poor comprehension, or uncooperativeness” (Heilman and Rothi, 1993). Apraxia is thus negatively defined, in terms of what it is not, as a higher-order disorder of movement that is not due to elementary sensory and/or motor deficits. This definition implies that there are situations where the effector is moved with normal skill (Hermsdörfer, Mai, Spatt, Marquardt, Veltkamp and Goldenberg, 1996). Puzzling parts of apraxia are the voluntary-automatic dissociation and context-dependence. On the one hand, apraxic patients may spontaneously perform gestures that they cannot perform on command (Schnider, Hanlon, Alexander and Benson, 1997). This voluntary-automatic dissociation can be illustrated by an apraxic patient that could use his left hand to shave and comb himself, but could not execute a specific motor action such as opening the
hand, so as to let go of an object (Lausberg, Göttet, Münssinger, Boegner and Marx, 1999). In this particular case, focussing on the target of the movement rather than on the movement itself increased his chances of a successful execution. On the other hand, the execution of the movement depends heavily on the context of testing (De Renzi, Faglioni and Sorgato, 1982). It may be well preserved in a natural context, with a deficit that appears in the clinical setting only, where the patient has to explicitly represent the content of the action outside of the situational props (Jeannerod and Decety, 1995; Leiguarda and Marsden, 2000).

Several authors agree that although apraxia is easy to demonstrate, it has proven difficult to understand. Research on apraxia is filled with confusing terminology, contradictory results and doubts that need to be resolved (Laeng, 2006; Goldenberg, Hermsdörfer and Spatt, 1996; De Renzi et al., 1982; Graham, Zeman, Young, Patterson and Hodges, 1999; Koski, Iacoboni and Mazziotta, 2002). Inconsistencies between similar studies may be explained by differences in methodological and statistical approaches in the apraxia assessment (i.e., types of gestures used and scoring criteria), chronicity and aetiology of damage and brain lesion localization tools (Haaland, Harrington and Knight, 2000). Therefore, it still stands that our understanding of the neural and cognitive systems underlying human praxis is not well established.

The chapter is structured as follows. We first review existing types of apraxia as well as important current and historical models of the apraxic deficit. We then consider the inter- and intra-hemispheric lesion correlates of apraxia. Two other sections are dedicated to the relationship between praxis and language and to the analysis of apraxic errors. We finally discuss the current state-of-art in apraxia, and argue for a multidisciplinary approach that encompasses evidence from various fields such as neuroimaging or neurophysiology.
Types of apraxia

This section reviews the current taxonomies of apraxia. Some of the frequently observed types of apraxia have inspired the apraxia models described in the following section, others still challenge them.

*Ideational apraxia* was historically defined as a disturbance in the conceptual organization of actions. It was first assessed by performing purposive sequences of actions that require the use of various objects in the correct order (e.g., preparing a cup of coffee) (Poeck, 1983). It was later accepted that ideational apraxia is not necessarily associated to complex actions, but is a larger deficit that also concerns the evocation of single actions. In this view, complex sequences of multiple objects are simply more suitable to reveal the deficit, possibly because of the heavier load placed on memory and attentional resources (De Renzi and Lucchelli, 1988). Nonetheless, the term *conceptual apraxia* was introduced to designate content errors in single actions, excluding sequence errors in multistaged actions with tools\(^1\) (Ochipa, Rothi and Heilman, 1992; Heilman, Maher, Greenwald and Rothi, 1997). In theoretical models, ideational and conceptual apraxia correspond to a disruption of the conceptual component of the praxis system, i.e., action semantics memory, described in more detail in the Models of apraxia section (De Renzi and Lucchelli, 1988; Graham et al., 1999). Patients with ideational apraxia are not impaired in the action execution *per se*, but demonstrate inappropriate use of objects and may fail in gesture discrimination and matching tasks. For example, a patient was reported to eat with a toothbrush and brush his teeth with a spoon and a comb. His inability to use tools could not be explained by a motor production deficit that would characterize ideomotor apraxia (defined below). Interestingly, although he was able to name the tools and point to them on command, he could not match the tools with the objects, hence suggesting a loss of knowledge related to the use of tools.

\(^1\)Conceptual apraxia is often observed in Alzheimer’s disease.
Ideomotor apraxia is considered to be a disorder of the production component of the praxis system, i.e., sensorimotor action programs that are concerned with the generation and control of motor activity (Rapcsak, Ochipa, Anderson and Poizner, 1995; Graham et al., 1999). It is characterized by errors in the timing, sequencing, and spatial organization of gestural movements (Leiguarda, 2001). Since the conceptual part of the praxis system is assumed to be intact, patients with ideomotor apraxia should not use objects and tools in a conceptually inappropriate fashion and should not have difficulty with the serial organization of an action (De Renzi et al., 1982). Ideational and ideomotor apraxia have been assessed by testing the execution of various types of gestures: transitive and intransitive (i.e., with or without the use of tools or objects), meaningless non-representational (e.g., hand postures relative to head) and meaningful representational (e.g., waving goodbye), complex sequences with multiple objects, repetitive movements, distal and proximal gestures (e.g., imitation of finger and hand configurations), reaching in peri-personal and body-centered space (e.g., targets in near space or on the patient’s body), novel movements (i.e., skill acquisition) or imagined movements. These gestures can also be executed under different modalities such as: verbal command, imitation, pantomime and tactile or visual presentation of objects.

The use of various gestures and different modalities to assess apraxia has helped to uncover many interesting functional dissociations that are listed below. For example, apraxia was shown to be modality-specific, i.e., the same type of gesture was differentially impaired according to the modality of testing (De Renzi et al., 1982). One dissociation, named conduction apraxia, is the syndrome of superior performance on verbal command than on imitation (Ochipa, Rothi and Heilman, 1994). The opposite pattern has also been observed: very poor performance on verbal command that improved on imitation or when seeing the object (Heilman, 1973; Merians, Clark, Poizner, Macauley, Gonzalez Rothi and Heilman, 1997). The extreme occurrence of conduction apraxia, namely the selective inability to imitate with normal performance on verbal command was termed visuo-imitative.
apraxia (Merians et al., 1997). In some cases of visuo-imitative apraxia, defective imitation of meaningless gestures (e.g., fist under chin) contrasts with preserved imitation of meaningful gestures (e.g., hitchhiking) (Goldenberg and Hagmann, 1997; Salter, Roy, Black, Joshi and Almeida, 2004). A surprising case of double dissociation from this kind of visuo-imitative apraxia was described in Bartolo, Cubelli, Sala, Drei and Marchetti (2001), where the patient showed impairment in meaningful gesture production (both on imitation and verbal command) and normal performance in imitation of meaningless gestures, suggesting that the patient was able to reproduce only movements he did not identify or recognize as familiar. Similarly, the apraxic patients in Buxbaum, Sirigu, Schwartz and Klatzky (2003) responded abnormally to familiar objects (e.g., a key, a hammer or a pen) but normally in recognizing the hand postures appropriate for novel objects (e.g., parallelepipeds differing in size and depth). These two studies argue that the reproduction of a gesture may be constrained by its degree of familiarity, indicating that current models of apraxia would need some refinement.

Furthermore, the representation of transitive and intransitive actions may be dissociable. In Watson, Fleet, Rothi and Heilman (1986), bilateral apraxia was observed only for transitive (e.g., hammering) but not intransitive (e.g., hitchhiking, waving goodbye) movements\(^2\). Whereas transitive gestures are constrained by the shape, size and function of objects, intransitive actions are related to socio-cultural contexts (Cubelli, Marchetti, Boscolo and Della Sala, 2000; Heath, Roy, Black and Westwood, 2001). The isolated disturbance of transitive hand movements for use of, recognition and interaction with an object, in the presence of preserved intransitive movements, was named tactile apraxia and usually appears in the hand contralateral to the lesion (Binkofski, Kunesch, Classen, Seitz and Freund, 2001).

As mentioned in the Introduction, contextual cues strongly influence the execution of actions. Some studies have systematically manipulated the contextual cues in order to

\(^2\)These patients had lesions in the left supplementary motor area (SMA).
assess their relative importance. For example, patients with impaired pantomime of motor actions showed no deficit in the comprehension of the use of tools or in manipulating the tools (Halsband, Schmitt, Weyers, Binkofski, Grützner and Freund, 2001). Graham et al. (1999) also observed dramatic facilitation in the demonstration of tool use when the patient was given the appropriate or a neutral tool to manipulate\(^3\). Interestingly, the patient could not prevent himself from performing the action appropriate to the tool he was holding, rather than the action that was requested. In another study however, gesture execution improved when the object of the action, but not the tool, was given (Clark, Merians, Kothari, Poizner, Macauley, Rothi and Heilman, 1994). Hence, the addition of visual and somaesthetic cues may improve certain aspects of apraxic movements, since it provides mechanical constraints and supplementary information that facilitates the selection of an adequate motor program (Hermsdörfer, Hentze and Goldenberg, 2006). Nonetheless, there is the case of a patient that performed much worse when he was actually manipulating the tool than on verbal command\(^4\) (Merians, Clark, Poizner, Jacobs, Adair, Macauley, Rothi and Heilman, 1999).

Dissociations that concern the nature of the target were also observed. For example, the left brain damaged patients in Hermsdörfer, Blankenfeld and Goldenberg (2003) had prolonged movement times and reduced maximum velocities when the movements were directed toward an allocentric target without visual feedback, but performed normally when the target was their own nose. Also, a clear dissociation was found in Ietswaart, Crey and Della Sala (2006) between impaired gesture imitation and intact motor programming of goal-directed movements, hence arguing against the interpretation of impaired imitation as a purely executional deficit (see the Models of apraxia section).

A particular type of apraxia is **constructional apraxia**, originally described by Kleist as “the inability to do a construction” and defined by Benton as “the impairment in combina-

\(^3\)The subject had clinically diagnosed corticobasal degeneration.

\(^4\)Ibid.
tor or organizing activity in which details must be clearly perceived and in which the relationship among the component parts of the entity must be apprehended" (Laeng, 2006). Constructional apraxic patients are unable to spontaneously draw objects, copy figures and build blocks or patterns with sticks, following damage to the dominant but also non-dominant hemisphere. Hence, constructional apraxia appears to reflect the loss of bilaterally distributed components for constructive planning and the perceptual processing of categorical and coordinate spatial relations (Platz and Mauritz, 1995; Laeng, 2006).

Apraxia can also be observed in mental motor imagery tasks. Motor imagery is considered as a means of accessing the mechanisms of action preparation and imitation, by sharing a common neural basis (Jeannerod and Decety, 1995). Apraxic patients were deficient in simulating hand actions mentally and in imagining the temporal properties of movements (Sirigu, Daprati, Pradat-Diehl, Franck and Jeannerod, 1999). Other apraxic patients showed a deficit in generating and maintaining internal models for planning object-related actions (Buxbaum, Johnson-Frey and Bartlett-Williams, 2005). These findings support the notion that the motor impairments observed in apraxic patients result from a specific alteration in their ability to mentally evoke actions, or to use stored motor representations for forming mental images of actions.

Apraxia may also be appropriate to reveal the role of feedback during the execution of a movement. Some apraxic patients were impaired in reaching and aiming movements only in the condition without visual feedback (Ietswaart, Crey, Della Sala and Dijkhuizen, 2001; Ietswaart et al., 2006) and performed worse during pointing with closed eyes (Hermsdörfer et al., 2003; Jacobs, Adair, Macauley, Gold, Gonzalez Rothi and Heilman, 1999). Interestingly, the patients in Haaland, Harrington and Knight (1999) overshot the target when feedback of the hand was removed and undershot the target when the feedback of the target was unavailable. Importantly, these patients continued to rely on visual feedback during the secondary adjustment phase of the movement and never achieved nor-

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5 These patients had posterior parietal lesions.
mal end-point accuracy when visual feedback of the hand position or target location was unavailable. These findings also suggest that ideomotor limb apraxia may be associated with the disruption of the neural representations for the extrapersonal (spatial location) and intrapersonal (hand position) features of movement (Haaland et al., 1999).

The importance of feedback signals was demonstrated in one of our own apraxic patients (unpublished data). We reproduced a seminal study of imitation of meaningless gestures by Goldenberg, Laimgruber and Hermsdörfer (2001) on an apraxic patient with left parietal ischemic lesion. We observed that the patient relied heavily on visual and tactile feedback. He often needed to bring his hand in the field of vision and corrected the hand posture by directly comparing it with the displayed stimulus to imitate. He also used tactile exploration when searching for the correct spatial position on his face. He showed many hesitations and extensive searching which led to highly disturbed kinematic profiles of the gesture (shown in Figure 4c, d), but often correct final postures.

Apraxia can also be defined in relation to the selectively affected effector: orofacial apraxia or buccofacial apraxia, oral apraxia, upper and lower face apraxia (Sala, Maistrello, Motto and Spinnler, 2006), lid apraxia, limb apraxia, leg apraxia, trunk apraxia, etc. Oral apraxia for example, is defined as the inability to perform mouth actions such as sucking from a straw or blowing a kiss. It should not be confounded with apraxia of speech (also called verbal apraxia), which is a selective disturbance of the articulation of words (Bizzozero, Costato, Sala, Papagno, Spinnler and Venneri, 2000). Motor planning disorders in children are denominated developmental dyspraxia (Cermak, 1985). Apraxia can also designate a praxic ability impaired in an isolated manner such as: gait apraxia, apraxic agraphia, dressing apraxia, orienting apraxia and mirror apraxia (i.e., inability to reach to objects in a mirror (Binkofski, Butler, Buccino, Heide, Fink, Freund and Seitz, 2003)). When the side of brain lesion and affected hand are considered, the terms sympathetic and crossed apraxia are used. Apraxia can sometimes be related to the spe-

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6Hand postures relative to the head, an example is shown in Figure 4a.
cific neural substrate that causes the disorder, for example following subcortical lesions in corticobasal degeneration (Pramstaller and Marsden, 1996; Jacobs et al., 1999; Merians et al., 1999; Hanna-Pladdy, Heilman and Foundas, 2001; Leiguarda, 2001) or following lesions of the corpus callosum (Watson and Heilman, 1983; Goldenberg et al., 2001; Lausberg et al., 1999; Lausberg, Davis and Rothenhäsler, 2000; Lausberg and Cruz, 2004). *Callosal apraxia* for example is particularly appropriate for disentangling the specific hemispheric contributions to praxis.

An extensive list of the types of apraxia and their definitions, including types that were not mentioned above, can be found in the Table in Figure 1.

**Models of apraxia**

Contemporary neuropsychological views of apraxia arise from Liepmann’s influential work that dates from more than a hundred years ago. Liepmann proposed the existence of an idea of the movement, “movement formulae”, that contains the “time-space-form picture” of the action (Rothi, Ochipa and Heilman, 1991). He believed that in right-handers, these movement formulae are stored in the left parietal lobe, endorsing the view of a left hemispheric dominance for praxis (Faglioni and Basso, 1985; Leiguarda and Marsden, 2000). To execute a movement, the spatio-temporal image of the movement is transformed into “innervatory patterns” that yield “positioning of the limbs according to directional ideas” (Jacobs et al., 1999). Liepmann distinguished between three types of apraxia, that correspond to disruptions of specific components of the model (Goldenberg, 2003; Faglioni and Basso, 1985). First, a damaged movement formula (i.e., faulty integration of the elements of an action) would characterize *ideational apraxia*. Second, failure of the transition from the movement formula to motor innervation (i.e., inability to translate a correct idea of the movement into a correct act) is defined as *ideomotor apraxia*. According to Liepmann, faulty imitation of movements is a purely executional deficit and proves the separation
between the idea and execution of a movement, since in imitation the movement formula is defined by the demonstration (Goldenberg, 1995; Goldenberg and Hagmann, 1997; Goldenberg, 2003). Finally, loss of purely kinematic (kinaesthetic or innervatory) inherent memories of an extremity is the limb-kinetic variant of apraxia.

Another historically influential model is the disconnection model of apraxia proposed by Geschwind (1965). According to this model the verbal command for the movement is comprehended in Wernicke's area and is transferred to the ipsilateral motor and premotor areas that control the movement of the right hand (Clark et al., 1994; Leiguarda and Marsden, 2000). For a left hand movement, the information needs to be further transmitted to the right association cortex via the corpus callosum. The model postulates that the apraxic disorder follows from a lesion in the left and right motor association cortices, or a disruption in their communication pathways. However this model cannot explain impaired imitation and impaired object use since these tasks do not require a verbal command (Rothi et al., 1991).

Heilman and Rothi (1993) proposed an alternative representational model of apraxia, according to which apraxia is a gesture production deficit that may result from the destruction of the spatiotemporal representations of learned movements stored in the left inferior parietal lobule. They proposed to distinguish between dysfunction caused by destruction of the parietal areas (where the spatiotemporal representations of movements would be encoded), and the deficit which would result from the disconnection of these parietal areas from the frontal motor areas (Heilman, Rothi and Valenstein, 1982). In the first case, posterior lesions would cause a degraded memory trace of the movement and patients would not be able to correctly recognize and discriminate gestures. In the second case, anterior lesions or disconnections would only provoke a memory egress disorder. Therefore patients with a gesture production deficit with anterior and posterior lesions should perform differently on tasks of gesture discrimination, gesture recognition, and novel gesture learning.
Roy and Square (1985) proposed a cognitive model of limb praxis that involves two systems, i.e., a conceptual and a production system (illustrated in Figure 2). The conceptual system provides an abstract representation of the action and comprises three kinds of knowledge: (1) knowledge of the functions of tools and objects, (2) knowledge of actions independent of tools and objects and (3) knowledge about the organization of single actions into sequences. The production system incorporates a sensorimotor representation of the action and mechanisms for movement control. Empirical support for the division of the praxis system into a conceptual and a production component is provided by a patient that could comprehend and discriminate transitive gestures she was unable to perform (Rapcsak et al., 1995). This model predicts three patterns of impairment (Heath et al., 2001). First, a deficit in pantomime but not in imitation would reflect damage to the selection and/or evocation of actions from long-term memory. Second, a deficit in imitation alone would indicate a disruption of the visual gestural analysis or translation of visual information into movement. Finally, concurrent impairment in pantomime and imitation is thought to reflect a disturbance at the latter, executive stage of gesture production and was the most frequent pattern observed in Roy, Heath, Westwood, Schweizer, Dixon, Black, Kalbfleisch, Barbour and Square (2000) and Parakh, Roy, Koo and Black (2004).

None of these models predict a number of modality-specific dissociations observed in neurologically impaired patients, such as preserved gesture execution on verbal command that is impaired in the visual modality when imitating (Ochipa et al., 1994; Goldenberg and Hagmann, 1997). To account for these dissociations, Rothi et al. (1991) proposed a cognitive neuropsychological model of limb praxis which reflects more appropriately the complexity of human praxis (illustrated in Figure 3a). This multi-modular model has input that is selective according to the modality, a specific “action semantics system” dissociable from other semantics systems, an “action reception lexicon” that communicates with an “action production lexicon” and a separate “nonlexical route” for the imitation of novel and
meaningless gestures\(^7\) (Rothi, Ochipa and Heilman, 1997).

Although this model is widely used to explain data from multiple neurological studies, it has difficulties concerning several aspects. First, it does not consider the existence of a selective tactile route to transitive actions (Graham et al., 1999). For example, the model fails to explain data from a patient profoundly impaired in gesturing in the verbal and visual modalities, but not with the tool in hand (Buxbaum, Giovannetti and Libon, 2000). Second, imitation of meaningless gestures is assumed to test the integrity of a direct route from visual perception to motor control. However, Goldenberg et al. (1996) have shown that this route is far from direct and involves complex intermediate processing steps. For example, apraxic patients that are impaired in reproducing gestures on their own bodies are also impaired in replicating the gestures on a life-sized mannikin (Goldenberg, 1995). Hence, general conceptual knowledge about the human body and the spatial configuration of body parts seems necessary for performing an imitation task (Goldenberg, 1995; Goldenberg et al., 1996; Goldenberg and Hagmann, 1997). The belief that imitation is a rather simple and straightforward visuomotor process is misleading as one would have to resolve the \textit{“body correspondence problem”}\(^8\) to transpose movements from bodies with different sizes and different owners which are in addition represented in different perspectives (Goldenberg, 1995).

To account for the last observation, Cubelli et al. (2000) have revised Rothi et al.’s cognitive neuropsychological model of limb praxis (illustrated in Figure 3b). They have added “a visuomotor conversion mechanism” devoted to transcoding the visual input into appropriate motor programs. They have also suppressed the direct link between the “input” and “output action lexicon”, leaving only an indirect link through the “action semantics

\(^7\)The vocabulary was borrowed from the literature of language processing.

\(^8\)Here we give a shortened version of the informal statement of the body correspondence problem. Given an observed behavior of the model, i.e., a sequence (or hierarchy) of subgoals, find and execute a sequence of actions using one own’s (possibly dissimilar) embodiment which leads through the corresponding subgoals (Nehaniv and Dautenhahn, 2002).
system”, as no empirical evidence was found of a patient able to reproduce familiar gestures with obscure meaning, but not unfamiliar gestures (see Figure 3a, b). Finally, they have also added a “gestural buffer” aimed at holding a short-term representation of the whole action. The model predicts five different clinical pictures (for definitions of the different apraxic disorders please refer to the Table in Figure 1): (1) a deficit of the “action input lexicon”: pantomime agnosia (i.e., a difficulty in the discrimination and comprehension of gestures) (2) a deficit of the “action semantics system”: conceptual apraxia without ideomotor apraxia, (3) a deficit of the “action output lexicon”: conceptual apraxia with spared gesture-meaning association, (4) a deficit of the “visuomotor conversion mechanism”: conduction apraxia (not observed in their study) and (5) a deficit of the “gestural buffer”: both ideomotor and ideational apraxia (i.e., impairment in all execution tasks with preserved ability to perform judgement and categorization tasks).

Buxbaum et al. (2000) further extended Rothi et al.’s cognitive neuropsychological model of limb praxis, based on their observation of a patient that performed particularly poorly on tasks that required a spatial transformation of the body. According to their model (illustrated in Figure 3c), a unitary set of representations named “body schema” calculates and updates the dynamic positions of the body parts relative to one another. Importantly, this dynamic body-centered representation of actions is a common processing stage between the “lexical” and “nonlexical route” and hence subserves both meaningful and meaningless actions. Note that at the level of the “lexical route”, there is an additional interaction with the stored representations of learned actions.

Existing models of apraxia still fail to account for additional empirical evidence such as for example, the differential performance in imitation of hand postures and imitation of finger configurations shown in Goldenberg and Hagmann (1997) and Goldenberg and Karnath (2006). Furthermore, in a study of ideomotor apraxia, Buxbaum, Kyle and Menon (2005) provided data which is compatible with the influential “mirror neuron hypothesis”. Apraxia models cannot easily be reconciled with this hypothesis which, based upon neu-
rophysiological observations from the monkey brain, postulates a “mirror neuron system” underlying both action recognition and action execution (Rizzolatti and Craighero, 2004). Mirror neurons are a special class of visuomotor neurons, initially discovered in area F5 of the monkey premotor cortex (see Figure 5), that discharge both when the monkey does a particular action and when it observes another individual doing a similar action (Gallese, Fadiga, Fogassi and Rizzolatti, 1996; Rizzolatti and Luppino, 2001; Rizzolatti, Fogassi and Gallese, 2002). Hence, the “mirror neuron system” is believed to map observed actions onto the same neural substrate used to execute these actions. As the same representations appear to subserve both action recognition and action production tasks, it would not be surprising if the perception of a movement is constrained by its executional knowledge. Related to apraxia, the “mirror neuron hypothesis” questions the separation of the “input” and “output lexicon” (Koski et al., 2002).

Contributions of the left and right brain hemispheres

Although most apraxia studies show a left brain hemisphere dominance for praxis, the studies arguing for a significant involvement of the right hemisphere are numerous. Left brain damage usually affects both hands, whereas right brain damage affects only the left hand, suggesting that the left hemisphere is fully competent for processing movement concepts and also contributes to the generation of movements in the right hemisphere. Apraxic deficits following left hemisphere lesions are also more frequent (De Renzi, Motti and Nichelli, 1980; Weiss, Dohle, Binkofski, Schnitzler, Freund and Heftner, 2001), however, in some rare cases, severe apraxia was observed following right hemisphere lesions (Marchetti and Sala, 1997; Raymer, Merians, Adair, Schwartz, Williamson, Rothi, Poizner and Heilman, 1999). The concept of crossed apraxia was introduced to describe patients with this opposite pattern of limb apraxia that cannot be explained by handedness. Callosal lesions
are most suitable for investigating the issues of hemispheric specialization of praxis. For example, split-brain patients were apraxic with their left hands, also suggesting a left hemisphere dominance for processing skilled movement (Watson and Heilman, 1983; Lausberg et al., 1999; Lausberg, Cruz, Kita, Zaidel and Ptito, 2003), but both hemispheres appeared to contain concepts for skill acquisition (Lausberg et al., 1999) and object use (Lausberg et al., 2003).

In kinematic studies (described in more detail in The analysis of apraxic errors section), only left brain damaged patients were impaired in imitation of meaningless movements (Hermsdörfer et al., 1996; Weiss et al., 2001), as well as in pointing movements (Hermsdörfer et al., 2003); whereas right brain damaged patients had deficits in slow-paced tapping and initiation of aiming movements (Haaland and Harrington, 1996). Hence, the left hemisphere was associated to movement trajectory control (Haaland, Prestopnik, Knight and Lee, 2004), sequencing and ballistic movements (Hermsdörfer et al., 2003) and the right hemisphere was related to on-line control of the movement (Hermsdörfer et al., 2003) and closed-loop processing (Haaland and Harrington, 1996).

A left-right dichotomy was also observed for imitation and matching of hand and finger configurations (Goldenberg, 1999). Left brain damaged patients had more difficulties with imitation than matching and vice-versa. In addition, the left hemisphere seemed fully competent for processing hand postures, but needed the right hemisphere’s contribution for processing finger postures (Goldenberg et al., 2001; Goldenberg, 2001; Sala, Faglioni, Motto and Spinnler, 2006). It was concluded that the left hemisphere mediates conceptual knowledge about the structure of the human body and that the right hemisphere is specialized for visually analyzing the gesture (Goldenberg, 1999; Goldenberg et al., 2001; Goldenberg, 2001).

Finally, several studies observed similar impairment scores following left and right brain lesions, arguing for a bihemispheric representation of skilled movement (Haaland and Flaherty, 1984; Kertesz and Ferro, 1984; Roy, Black, Winchester and Barbour, 1992; Roy
et al., 2000; Heath et al., 2001). The less frequent, nevertheless well detected incidence of limb apraxia following right brain lesion, was attributed to the sensitivity and precision of the assessment methodology. In addition, right hemisphere lesions often led to severe face apraxia (Bizzozero et al., 2000; Sala, Maistrello, Motto and Spinnler, 2006). Hence, a model of widespread praxis, distributed across both hemispheres, may be more appropriate than the unique left lateralised center previously hypothesized. Moreover, it seems that the degree of left hemisphere dominance varies within subjects and with the type of movement (Haaland et al., 2004), raising the issue of overlap between the contributions of the right and left hemisphere to specialized praxic functions.

Intrahemispheric lesion location: a distributed representation of praxis?

Several studies have failed to find a consistent association between the locus of the lesion within a hemisphere and the severity of apraxia (Basso, Luzzatti and Spinnler, 1980; Kertesz and Ferro, 1984; Alexander, Baker, Naeser, Kaplan and Palumbo, 1992; Schnider et al., 1997; Hermsdörfer et al., 2003). Moreover, areas involved in apraxia can also be damaged in non-apraxic patients (Haaland et al., 1999; Buxbaum et al., 2003). However, apraxic deficits are most frequent following parietal and frontal lesions, but were also observed in patients with temporal, occipital and subcortical damage (De Renzi and Lucchelli, 1988; Goldenberg, 1995; Hermsdörfer et al., 1996; Bizzozero et al., 2000).

More specifically, ideomotor apraxia and motor imagery deficits were observed following lesions in the left inferior parietal and the left dorsolateral frontal lobes (Haaland et al., 2000; Buxbaum, Johnson-Frey and Bartlett-Williams, 2005). For example, several studies suggested that Brodmann areas 39 and 40 (i.e., angular and supramarginal gyri of the inferior parietal lobule) are critical in visuo-imitative apraxia (Goldenberg and Hagmann,
and ideomotor limb apraxia (Haaland et al., 1999; Buxbaum et al., 2003). In addition, the superior parietal lobe appeared crucial in integrating external visual and intrapersonal somaesthetic information (Heilman, Rothi, Mack, Feinberg and Watson, 1986; Haaland et al., 1999). Goldenberg and Karnath (2006) subtracted the lesion overlay of unimpaired from impaired patients and associated disturbed imitation of hand postures with lesions in the inferior parietal lobe and temporo-parieto-occipital junction, whereas disturbed imitation of finger postures could be related to lesions in the inferior frontal gyrus. Interestingly, parts of the middle and inferior frontal gyri, in the vicinity of Brodmann areas 6, 8 and 46, were involved in all of the ideomotor apraxics in Haaland et al. (1999). Furthermore, premotor lesions (including lesions to the supplementary motor area) particularly affected bimanual actions in Halsband et al. (2001) and transitive actions in (Watson et al., 1986).

It has been difficult to disentangle between the specific contributions of the parietal and the frontal cortices, as lesions in these areas lead to similar deficits (Haaland et al., 1999; Haaland et al., 2000). For example, target and spatial errors were related to posterior lesions only (Haaland et al., 2000; Halsband et al., 2001; Weiss et al., 2001; Goldenberg and Karnath, 2006), but internal hand configuration errors were present in patients with anterior and posterior lesions (Haaland et al., 2000; Goldenberg and Karnath, 2006). Importantly, only patients with posterior lesions, and not anterior lesions, had difficulties with discriminating between correctly and incorrectly performed actions and with recognizing pantomimes or appropriate hand postures (Halsband et al., 2001; Buxbaum, Kyle and Menon, 2005).

Apraxia can also develop following subcortical lesions (Pramstaller and Marsden, 1996; Graham et al., 1999; Jacobs et al., 1999; Merians et al., 1999; Hanna-Pladdy et al., 2001). In this case, it is not clear whether the apraxia originates from lesions in the basal ganglia, which are extensively connected to the superior parietal lobe and premotor and supplementary motor areas (Jacobs et al., 1999; Merians et al., 1999), or from the surrounding
white matter (i.e., frontoparietal connections) (Pramstaller and Marsden, 1996).

Failure to find clear correlations between specific lesion loci and different apraxic deficits argues for a wide-spread cortical and subcortical representation of praxis, distributed across specialized neural systems working in concert (Hermsdörfer et al., 2003; Leiguarda and Marsden, 2000). However, we believe that a selective damage to one of these systems may produce a particular pattern of errors tightly related to a subtype of apraxia.

Praxis and Language?

Apraxia is most often seen in association with aphasia (i.e., loss of the ability to speak or understand speech), which renders the assessment of apraxia very difficult. Indeed, one has to provide evidence that the patient has understood the commands so that the motor deficit cannot be attributed to aphasia (De Renzi et al., 1980). Historically, gestural disturbance in aphasics was considered to be a manifestation of damaged abstract knowledge. This idea of a common impaired symbolic function underlying aphasia and apraxia was supported for a long time (Kertesz and Hooper, 1982). However, several large-scale studies failed to find correlations between subtypes of apraxia and aphasia (Goodglass and Kaplan, 1963; Lehmkuhl, Poeck and Willmes, 1983; Buxbaum, Kyle and Menon, 2005). Moreover, clear evidence of a double dissociation between apraxia and aphasia was presented in Papagno, Della Sala and Basso (1993). For example, some patients were able to verbalize a desired movement but could not perform it (Goodglass and Kaplan, 1963), whereas other patients were able to pantomime actions they were unable to name (Rothi et al., 1991). Hence, it seems that many aspects of language and praxis are subserved by independent, possibly contiguous neuronal processes, but concomitant deficits may also appear because of shared neuroanatomical substrates (Kertesz and Hooper, 1982). Nevertheless, the question of how language is related to praxis is a fascinating one and needs further study, as it can give some insight into the existence of a supramodal representation of knowledge, or alternatively
shed light onto the communication mechanisms between the praxic- and language- specific representations of knowledge\textsuperscript{9}.

**The analysis of apraxic errors**

There are extensive quantitative analyses of the severity of apraxic errors in single case studies and in large samples of brain damaged patients. *Qualitative analyses* however are less numerous and unstandardized, but nonetheless essential for precisely understanding the nature of apraxia. Performances are usually classified in a limited number of response categories such as\textsuperscript{10}: temporal errors, spatial errors, content errors, substitutive errors, augmentative errors, fragmentary errors, associative errors (i.e., the correct movement is replaced by another movement that shares one feature), parapraxic errors (i.e., correct execution of a wrong movement), wrong body part errors (e.g., patients that execute a correct movement with the leg instead of the hand), body part as tool errors (i.e., a body part is used to represent the imagined tool) and perseveration errors (Lehmkuhl et al., 1983; Platz and Mauritz, 1995; Poeck, 1983; De Renzi and Lucchelli, 1988; Halsband et al., 2001; Weiss et al., 2001; Lausberg et al., 1999; Lausberg et al., 2003). Perseveration and body parts as tool errors should be accorded some special interest in future studies, as they are prominent in apraxia and their occurrence is far from being elucidated (Poeck, 1983; Raymer, Maher, Foundas, Heilman and Rothi, 1997; Lausberg et al., 2003). For example, even though normal subjects also commit body part as tool errors\textsuperscript{11}, only subjects with brain lesion cannot correct their error after reinstruction (Raymer et al., 1997).

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\textsuperscript{9}Some authors have posited that an action-recognition mechanism might be at the basis of language development (Rizzolatti and Arbib, 1998).

\textsuperscript{10}This list is not extensive. Terminologies can vary a lot across different authors.

\textsuperscript{11}There is a hierarchical organization in the performance of actions with increasing difficulty. Children first acquire the ability to actually use objects, then to demonstrate the action with similar substitute objects, then with dissimilar substitute objects, then to use body parts as substitutes, and finally to perform pantomimes with holding imagined objects. This note was taken from Lausberg et al. (2003).
A significant step forward in the analysis of apraxic errors was the use of quantitative 3D kinematic motion analysis. These techniques allowed to show many abnormalities in the kinematic features of apraxic movements such as for example: deficits in spatial accuracy, irregular velocity profiles, reduced maximum velocities, reduced movement amplitudes, decoupling of the relationship between instantaneous wrist velocity and trajectory curvature, improper linearity of the movement, wrong orientation of the movement in space and/or deficient joint coordination (Poizner, Mack, Verfaellie, Rothi and Heilman, 1990; Platz and Mauritz, 1995; Rapcsak et al., 1995; Poizner, Clark, Merians, Macauley, Rothi and Heilman, 1995; Poizner, Merians, Clark, Rothi and Heilman, 1997; Merians et al., 1999; Clark et al., 1994; Haaland et al., 1999; Binkofski et al., 2001; Hermsdörfer et al., 2006). An example of an apraxic movement with abnormal kinematics is shown in Figure 4. Based on kinematic studies it could be concluded that ideomotor limb apraxia impaired the response implementation but not the preprogramming of the movement (Haaland et al., 1999) and decoupled the spatial and temporal representations of the movement (Poizner et al., 1990; Poizner et al., 1995). Importantly, the kinematic abnormalities observed were often spatial and not temporal, the longer movement times in the apraxic group could be interpreted as an artefact of the longer distance traveled (Haaland et al., 1999; Hermsdörfer et al., 2006). However, several authors have advised against systematically interpreting the irregular kinematics as an indicator for deficient motor programming or deficient motor implementation (Platz and Mauritz, 1995; Haaland et al., 1999). For example, no correlation could be found between the kinematic abnormalities and apraxic errors in Hermsdörfer et al. (1996). Indeed, movements with degraded kinematics frequently reached a correct final position, while, on the contrary, kinematically normal movements often led to apraxic errors. The abnormal kinematic profile of the gesture probably arose from several corrective and compensatory strategies that the patient used to cope with the apraxic deficit (Hermsdörfer et al., 1996; Goldenberg et al., 1996). For example, hesitant and on-line controlled movements generated multi-peaked velocity
profiles in our study (see Figure 4d). Hence, according to the authors, the basic deficit underlying apraxia may concern the mental representation of the target position. Consistently with this hypothesis, it was found that apraxic patients relied more than normal subjects on online visual information in aiming movements (Ietwaart et al., 2006). Figure 4

Discussion

We have shown in the preceding sections that apraxia has proven very difficult to assess and understand. Here we will try to provide some hypotheses why these difficulties might arise and we propose several ways to overcome these.

The complex nature of apraxia. Apraxia designates the impairment of the human praxis system following brain lesion and has to deal with the high complexity and wide range of human praxic functions. Therefore studies of apraxia have separately tackled the faulty execution of many types of gestures (e.g., transitive and intransitive, meaningful and meaningless, peripersonal and body-centered, etc..) of various end-effectors (e.g., mouth, face, leg, limb) in different types of modalities (e.g., visual, auditive, tactile presentation and imitation). The high dimensionality of varying parameters has led to a lack of systematicity in the apraxia assessment and terminologies used. This has also rendered the coherent interpretation of the disorder rather arduous.

It follows that there is a great need to discriminate between different types of actions, as they appear to be differentially impaired in apraxia and hence may involve distinct underlying mechanisms (see the Types of apraxia section). Indeed, it is very likely that the mechanisms of imitation and execution of movements vary according to the type of action that is imitated or executed (Schnider et al., 1997; Goldenberg, 1999; Goldenberg and Karnath, 2006). This suggests that different categories of actions require the use of separate systems at some stage of the processing, but the level of separation between the
representations underlying actions of different types, or even different actions of the same type, is not at all clear yet.

We will principally argue that it is important to better understand what a particular gesture or execution modality implies in terms of brain resources and brain processes when compared to another gesture/execution modality. For example, a transitive action, i.e., an action that involves an object, is very different from an intransitive action in the sense that it provides supplementary tactile input as a result from the interaction with the object. This tactile sensory input then needs to be integrated to the representation of the action that relies also on other types of sensory inputs such as visual and proprioceptive. Moreover, executing a transitive action in a pantomime condition is also different from executing it with the object in hand, since the action has to be retrieved without the help of tactile input produced by the object. Indeed the movement is somehow modified, for example movement amplitudes in normal subjects were larger in the pantomime condition when compared to actual sawing (Hermsdörfer et al., 2006).

The distinction between meaningful and meaningless gestures would also need some clarification. The reproduction of a recognized meaningful gesture on the one hand, appears entirely based on the internal representation of the gesture. Indeed, the knowledge of a learned skilled act is preferably retrieved from motor memory rather than being constructed de novo (Halsband et al., 2001). On the other hand, the reproduction of a meaningless gesture involves a close visual tracking of the imitatee’s body configuration and was modeled by a “visuo-motor conversion mechanism” or a “body schema” (see Figure 3b, c). To summarize, a meaningful gesture seems to be, to a certain extent, assimilated to a goal that guides the action from memory, whereas a meaningless gesture is defined as a particular configuration of the body in space and time, with no external referents (Goldenberg, 2001). Hence, imitation of meaningless gestures might be used to test the comprehension and replication of changing relationships between the multiple parts and subdivisions of the refined and complex mechanical device which is the human body (Goldenberg, 2001). Further-
more, a preserved imitation of meaningless gestures is crucial for the apraxic patient as it might be useful for relearning motor skills. The double dissociation observed between imitation of meaningless and meaningful gestures argues for completely separate processing systems and is still not accounted for by any of the existing apraxia models previously described. However, meaningless actions involve novel motor sequences that must be analyzed and constructed from existing movements (Koski et al., 2002) and both meaningless and meaningful gestures appear to involve a body schema, i.e., a dynamic model for coding the body (Buxbaum et al., 2000). Hence, meaningless and meaningful actions may also share some overlapping conceptual representations.

These examples show that there are some common and some distinct processes involved in the different types of movements and modalities used for testing apraxia. Identifying the overlap of these processes would provide a clearer framework for interpreting the patient’s performance and would simplify the analysis of the lesion correlates. The choice of the testing condition is crucial, as well as identifying the processes inherent to the chosen condition. However this is a difficult task, since correlations can be found between some very different and even dissociated types of movements\textsuperscript{12}. For example, kinematic measures of pointing movements were correlated to gesture imitation, suggesting that the kinematic deficits observed during pointing movements are generalized to more global aiming movements, including movements for imitating hand gestures (Hermsdörfer et al., 2003). Accordingly, gesture imitation is believed to depend upon some of the same cognitive mechanisms as reaching and grasping (Haaland et al., 2000), however the level and extent of interplay is not clear. To make the picture even more complex, the underlying representations may be componential, for example with separate hand posture representations for transitive gestures (Buxbaum, Kyle and Menon, 2005). This leads us to two questions that urge to be answered: (1) what are the basic motor primitives from which all movements are con-

\textsuperscript{12}Surprisingly, single finger tapping was a better predictor of the severity of apraxia than goal-directed grasping and aiming (Ietswaart et al., 2006). Single finger tapping is almost never used to assess apraxia.
structed and (2) which are the motor components that are related to specific movements.

**Beyond the complex nature of apraxia.** One way to cope with the complex nature of apraxia is to be even more *precise* and *systematic* in assessing the apraxic disorder. Ideally, the full range of praxic functions, related to different effectors, including mouth, face and foot should be tested in a complete set of modalities (Koski et al., 2002). Moreover, we find unfortunate that qualitative measures of the errors, such as kinematic measures of the movement trajectory (refer to the The analysis of apraxic errors section), are frequently missing or given in a purely statistical fashion (e.g., 25% of errors in condition A). As such, these measures do not suffice to understand why the patient succeeds at the execution of some actions, but not other similar actions. For example, in one study the patient was able to evoke some actions (using a razor and a comb) fairly consistently, yet others (hammering and writing) were never produced (Graham et al., 1999). In another study, the same gestures were not always congruently disturbed across the different modes of execution, namely on imitation and on verbal command (Jacob s et al., 1999). We believe that it is this inability to *distinguish between different types of errors related to different types of gestures* that has prevented us so far from discovering the precise neuroanatomical correlates of apraxia, on top of the difficulty to accurately identify the brain lesion. Hence, the typology and analysis of apraxic errors need to be improved. We encourage extensive categorization of the errors and their characterization via kinematic methods. In addition, the errors should be reported *in relation to* the exact movement and not only specific condition tested.

We also suggest that studies that assess apraxia should more often integrate tasks of *motor learning*, as patients with apraxia may also be deficient in learning new motor tasks (Heilman, Schwartz and Geschwind, 1975; Rothi and Heilman, 1984; Platz and Mauritz, 1995; Lausberg et al., 1999). The main motivation in understanding apraxia is to help the apraxic patients in their everyday lives through the development of efficient
rehabilitation methods and training programmes\textsuperscript{13}. Assessing the exact expression of the apraxic deficit and especially the patient’s motor learning abilities, would help to choose an appropriate therapy for the patient. Efficiently targeting the movements and praxis components specifically affected in each patient would accelerate the process of improving his or her praxic faculties. For the moment, apraxia in relation to motor learning is an underinvestigated line of research.

Furthermore, we believe that \textit{modeling research} may prove very helpful to gain some insight into the details and potential implementation of the processes underlying human praxis. When a roboticist searches for an algorithm for his robot to manipulate objects, he or she has to provide with all the different input signals and implement in practice all the necessary computations and processing resources. For example, the differences and similarities between reaching to body-centered versus peripersonal cues would become evident through the development of corresponding algorithms, as they would be explicitly computed. According to Schaal and Schweighofer (2005), computational models of motor control in humans and robots often provide solid foundations that can help us to ground the vast amount of neuroscientific data that is collected today. Thus, biologically inspired modeling studies such as Sauser and Billard (2006) and Hersch and Billard (2006) seem to be very promising approaches in the understanding of the nature of gestures and in emphasizing the differences and similarities of the underlying processes.

Although neuropsychological models are essential for the understanding of apraxia, they do not address the question of the precise neural representation of the action and how this representation can be accessed. In a \textit{neurocomputational model}, one has to take into account the computational principles of movement that reproduce the behavioral and kinematic results of the patient, as well as propose a biologically plausible implementation of the black-box components of apraxia models. In this view, we have developed a simple

\textsuperscript{13}According to Platz and Mauritz (1995), only patients with ideomotor apraxia and not ideational and constructional apraxia could benefit from a task-specific sensorimotor training.
neurocomputational model described in Petreska and Billard (2006), that accounts for the callosal apraxic deficit observed in a seminal experimental study of imitation of meaningless gestures (Goldenberg et al., 2001). Our model combines two computational methods for unsupervised learning applied to a series of artificial neural networks. The biologically inspired and distributed representations of sensory inputs self-organize according to Kohonen’s algorithm and associate with antihebbian learning. The appropriate transformations between sensory inputs needed to reproduce certain gestures are thus learned within a biologically plausible framework. It is also possible to impair the networks in a way that accounts for the performance of Goldenberg et al.’s apraxic patient in all of the conditions of the study. The model also suggests potential neuroanatomical substrates for this task. We believe that the development of neurocomputational models is a good way to probe our understanding of apraxia and is compatible with the view of integrating knowledge from different lines of research, a point which we will defend in the following section.

Toward a multidisciplinary approach. We believe that apraxia can be best dismantled by adopting a multidisciplinary approach. Future models of apraxia will need to encompass knowledge and data from studies of normal human motor control, human brain imaging and monkey brain neurophysiology. Fortunately, several authors have already attempted to combine different sources of evidence: by considering apraxia in the neurophysiological framework (e.g., Leiguarda and Marsden (2000)) or by validating a model of apraxia using neuroimaging methods (e.g., Hermsdörfer, Goldenberg, Wachsmuth, Conrad, Ceballos-Baumann, Bartenstein, Schwaiger and Boecker (2001), Peigneux, van der Linden, Garraux, Laureys, Degueldre, Aerts, Del Fiore, Moonen, Luxen and Salmon (2004), Chaminade, Meltzoff and Decety (2005), Mühlau, Hermsdörfer, Goldenberg, Wohlschlager, Castrop, Stahl, Röttinger, Erhard, Haslinger, Ceballos-Baumann, Conrad and Boecker (2005)).

Normal human motor control has been extensively studied via behavioral, psychophys-
ical, kinematic or computational methods for decades, giving rise to several well established *principles of movement*, such as: spatial control of arm movements (Morasso, 1981), maps of convergent force fields (Bizzi, Mussa-Ivaldi and Giszter, 1991), uncontrolled manifold concept (Scholz and Schöner, 1999), \( \tau \)-coupling in the perceptual guidance of movements (Lee, Craig and Grealy, 1999) and inverse and forward internal models (Wolpert and Ghahramani, 2000). Studies of motor control have also inspired several models for reaching like: minimum jerk trajectory control (Flash and Hogan, 1985), vector-integration-to-endpoint model (Bullock and Grossberg, 1988), minimum torque change model (Uno, Kawato and Suzuki, 1989) and stochastic optimal feedback control (Todorov and Jordan, 2002) (for a review refer to Desmurget, Pélisson, Rossetti and Prablanc (1998)). Proposed models for grasping (e.g., schema design (Oztop and Arbib, 2002)) are reviewed in Jeannerod, Arbib, Rizzolatti, and Sakata (1995) and models for sensorimotor learning such as the modular selection and identification for control model (Haruno, Wolpert and Kawato, 2001) in Wolpert, Ghahramani and Flanagan (2001). In addition, it was also shown that the amplitude and direction of pointing movements may be independently processed (Vindras, Desmurget and Viviani, 2005) or that the kinematics and dynamics for reaching may be separately learned (Krakauer, Ghilardi and Ghez, 1999). Investigation of apraxia can only benefit from taking into account the rich knowledge of the computational processes of movement used by the brain and obviously, apraxia models would need to be compatible with the current general theories of movement control.

Progress in describing the contribution of specific brain regions to human praxis through the study of brain-damaged patients has been limited by the variability in the size, location and structures affected by the lesion (Koski et al., 2002). *Human brain imaging studies*, particularly positron emission tomography (PET) and functional magnetic resonance (fMRI) overcome this difficulty to a certain extent and have an essential role in resolving the neuroanatomical correlates of human functions. Despite the evident difficulties and limitations to study movements with neuroimaging, numerous studies have addressed the question of
the representation of human praxis, making significant contributions to the understanding of the neural substrates underlying visuomotor control (see Culham, Cavina-Pratesi and Singhal (2006) for a review). In order to give an idea of the number of praxis functions that have been addressed with brain imaging technologies, we will mention some of them: observation of meaningful and meaningless actions with the intent to recognize or imitate (Decety, Grèzes, Costes, Jeannerod, Procyk, Grassi and Fazio, 1997), hand imitation (Krams, Rushworth, Deiber, Frackowiak and Passingham, 1998), visually guided reaching (Kertzman, Schwarz, Zeffiro and Hallett, 1997; Desmurget, Epstein, Turner, Prablanc, Alexander and Grafton, 1999; Grefkes, Ritzl, Zilles and Fink, 2004), object manipulation and tool-use (Binkofski, Buccino, Stephan, Rizzolati, Seitz and Freund, 1999; Johnson-Frey, Newman-Norlund and Grafton, 2005), real and/or imagined pantomimes (Moll, de Oliveira-Souza, J., Cimini Cunha, Souza-Lima and Andreiuolo, 2000; Choi, Na, Kang, Lee, Lee and Na, 2001; Rumiati, Weiss, Shallice, Ottoboni, Noth and Zilles, 2004) and sequential organization of actions (Ruby, Sirigu and Decety, 2002). The areas specialized for the perception of body parts and postures have been consistently identified\textsuperscript{14} (Peigneux, Salmon, van der Linden, Garraux, Aerts, Delfiore, Degueldre, Luxen, Orban and Franck, 2000; Downing, Jiang, Shuman and Kanwisher, 2001). Most importantly, several brain imaging studies have been conducted in relation to apraxia (Hermsdörfer et al., 2001; Peigneux et al., 2004; Chaminade et al., 2005; Mühlau et al., 2005) with the intent to test the neuroanatomical hypothesis of the neuropsychological models previously described.

\textit{Neurophysiological studies} allow the investigation of brain processes at the neuronal level and are essential to the understanding of the principles of neural computation. Certainly the monkey brain differs from the human brain, however this discrepancy can be overcome to some extent through the search of homologies (Orban, Van Essen and Van-

\textsuperscript{14}Interestingly, these occipital and visually specialized areas are not only modulated by the visual presentation of body configurations, but also when the person executes a limb movement (Astaipayev, Stanley, Shulman and Corbetta, 2004), indicating a bidirectional flow of the information.
duffel, 2004; Sereno and Tootell, 2005; Arbib and Bota, 2003; Rizzolatti et al., 2002).

Sensorimotor processes such as reaching and grasping for example, have been extensively studied: several parallel parietofrontal circuits were identified, each subserving a particular sensorimotor transformation (Kalaska, Scott, Cisek and Sergio, 1997; Wise, Boussaoud, Johnson and Caminiti, 1997; Matelli and Luppino, 2001; Battaglia-Mayer, Caminiti, Lacquiniti and Zago, 2003). Without going into the details of the representations used in each of these functionally distinct parietal and frontal areas (illustrated in Figure 5), we will mention those which seem relevant for models of apraxia. For example, LIP-FEF neurons discharge in relation with eye movements and are sensitive to the direction and amplitude of eye saccades (Platt and Glimcher, 1998), VIP-F4 neurons construct the ‘peripersonal’ space confined to the head (Duhamel, Colby and Goldberg, 1998), AIP-F5 neurons mediate motor responses selective for hand manipulation and grasping movements (Cohen and Andersen, 2002), MIP-F2 neurons have a crucial role in the planning, execution and monitoring of reaching movements (Simon, Mangin, Cohen, Le Bihan and Dehaene, 2002) where MIP neurons respond to joint rotation (Eskandar and Assad, 1999) and F2 neurons are selective for grip and wrist orientation (Raos, Umiltá, Gallese and Fogassi, 2004). Furthermore, multiple space representations appear to coexist in the brain that integrate multisensory inputs (e.g., visual, somatosensory, auditory and vestibular inputs) (Graziano and Gross, 1998). For example, neurons in area 5 appear to combine visual and somatosensory signals in order to monitor the configuration of the limbs (Graziano, Cooke and Taylor, 2000) and the receptive fields of VIP neurons respond congruently (i.e., with matching receptive fields) to tactile and visual stimulation (Duhamel et al., 1998). It is very interesting that the modality-specific activities are spatially aligned: the visual receptive field corresponding to the arm or the face may shift along with that body part when it is passively moved (Graziano, Hu and Gross, 1997). In addition, neurophysiological data can give us insight into how the arm posture modulates the activity of somatosensory neurons (Helms Tillery, Soechting and Ebner, 1996) and how it affects the neurons that compute the
trajectory of the hand (Scott, Sergio and Kalaska, 1997). It should be noted that several sensorimotor transformations are needed in order to grasp an object, the motor command being in hand coordinates and the object’s location in gaze coordinates. To compute these transformations, the brain appears to use *multiple body-centered frames of references* (Graziano and Gross, 1998): the frames of references underlying VIP area neurons appear to be organized along a continuum from eye to head coordinates (Duhamel, Bremner, Ben-Hamed and Graf, 1997; Avillac, Denève, Olivier, Pouget and Duhamel, 2005) and direct transformations from head to body-centered representations are possible in the posterior parietal cortex (Buneo, Jarvis, Batista and Andersen, 2002; Buneo and Andersen, 2006) with an error estimate of the target position computed in a common eye reference frame (Batista, Buneo, Snyder and Andersen, 2002; Cohen and Andersen, 2002). Finally, it was also shown that tools may be integrated into the “body schema” at the neuronal level (Iriki, Tanaka and Iwamura, 1996; Maravita, Spence and Driver, 2003).

To conclude, we strongly believe that this multidisciplinary approach should be bidirectional. Not only apraxia can be interpreted in the neuropsychological and neurophysiological frameworks, but these research domains would also benefit from taking into consideration observations from apraxia. For example, one could learn enormously on how the normal human praxis system functions by looking at how it is affected by apraxia.

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References


<table>
<thead>
<tr>
<th>Type of Apraxia</th>
<th>Definition</th>
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</thead>
<tbody>
<tr>
<td>Ideational apraxia</td>
<td>Initially used to refer to impairment in the conceptual organization of actions, assessed with sequential use of multiple objects. Later defined as conceptual apraxia.</td>
</tr>
<tr>
<td>Conceptual apraxia</td>
<td>Impairment in the concept of a single action, characterized by content errors and the inability to use tools.</td>
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<tr>
<td>Ideomotor apraxia</td>
<td>Impairment in the performance of skilled movements, characterized by spatial or temporal errors in the execution of movements.</td>
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<tr>
<td>Limb-kinetic apraxia</td>
<td>Slowness and stiffness of movements with a loss of fine, precise and independent movement of the fingers.</td>
</tr>
<tr>
<td>Constructional apraxia</td>
<td>Difficulty in drawing and constructing objects. Impairment in the combinatory or organizing activity in which details and relationship among the component parts of the entity must be clearly perceived.</td>
</tr>
<tr>
<td>Developmental dyspraxia</td>
<td>Disorders affecting the initiation, organization and performance of actions in children.</td>
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<tr>
<td>Modality-specific apraxias</td>
<td>(localized within one sensory system)</td>
</tr>
<tr>
<td>Pantomime agnosia</td>
<td>Normal performance in gesture production tests both on imitation and on verbal command, but poor performance in gesture discrimination and comprehension. Patients with pantomime agnosia can imitate pantomimes they can not recognize.</td>
</tr>
<tr>
<td>Conduction apraxia</td>
<td>Superior performance on pantomime to verbal command than on pantomime imitation.</td>
</tr>
<tr>
<td>Visuo-imitative apraxia</td>
<td>Normal performance on verbal command with selectively impaired imitation of gestures. Also used to designate the defective imitation of meaningless gestures combined with preserved imitation of meaningful gestures.</td>
</tr>
<tr>
<td>Optical (or visuomotor) apraxia</td>
<td>Disruptions to actions calling upon underlying visual support.</td>
</tr>
<tr>
<td>Tactile apraxia</td>
<td>Disturbance of transitive hand movements for use of, recognition and interaction with an object, in the presence of preserved intransitive movements.</td>
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<tr>
<td>Effector-specific apraxias</td>
<td></td>
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<tr>
<td>Upper/lower face apraxia</td>
<td>Impairment in performing actions with parts of the face.</td>
</tr>
<tr>
<td>Oral apraxia</td>
<td>Inability to perform skilled movements with the lips, cheeks and tongue.</td>
</tr>
<tr>
<td>Orofacial (or buccofacial) apraxia</td>
<td>Difficulties with performing intentional movements with facial structures including the cheeks, lips, tongue and eyebrows.</td>
</tr>
<tr>
<td>Lid apraxia</td>
<td>Difficulty with opening the eyelids.</td>
</tr>
<tr>
<td>Ocular apraxia</td>
<td>Impairment in performing saccadic eye movements on command.</td>
</tr>
<tr>
<td>Limb apraxia</td>
<td>Used to refer to ideomotor apraxia of the limbs frequently including the hands and fingers.</td>
</tr>
<tr>
<td>Trunk (or axial) apraxia</td>
<td>Difficulty with generating body postures.</td>
</tr>
<tr>
<td>Leg apraxia</td>
<td>Difficulty with performing intentional movements with the lower limbs.</td>
</tr>
<tr>
<td>Task-specific apraxias</td>
<td></td>
</tr>
<tr>
<td>Gait apraxia</td>
<td>Impaired ability to execute the highly practised, co-ordinated movements of the lower legs required for walking.</td>
</tr>
<tr>
<td>Gaze apraxia</td>
<td>Difficulty in directing gaze.</td>
</tr>
<tr>
<td>Apraxia of speech (or verbal apraxia)</td>
<td>Disturbances of word articulation.</td>
</tr>
<tr>
<td>Apraxic agraphia</td>
<td>A condition in which motor writing is impaired but limb praxis and nonmotor writing (typing, anagram letters) are preserved.</td>
</tr>
<tr>
<td>Dressing apraxia</td>
<td>Inability to perform the relatively complex task of dressing.</td>
</tr>
<tr>
<td>Dyssynchronous apraxia</td>
<td>Failure to combine simultaneous preprogrammed movements.</td>
</tr>
<tr>
<td>Orienting apraxia</td>
<td>Difficulty in orienting one’s body with reference to other objects.</td>
</tr>
<tr>
<td>Mirror apraxia</td>
<td>A deficit in reaching to objects presented in a mirror.</td>
</tr>
<tr>
<td>Lesion-specific apraxias</td>
<td></td>
</tr>
<tr>
<td>Callosal apraxia</td>
<td>Apraxia caused by damage to the anterior corpus callosum that usually affects the left limb.</td>
</tr>
<tr>
<td>Sympathetic apraxia</td>
<td>Apraxia of the left limb due to damage to the anterior left hemisphere (the right hand being partially or fully paralysed).</td>
</tr>
<tr>
<td>Crossed apraxia</td>
<td>The unexpected pattern of apraxia of the right limb following damage to the right-hemisphere.</td>
</tr>
</tbody>
</table>

Figure 1: Taxonomy of apraxia.
Conceptual System

Abstract knowledge of Action:
- Knowledge of Object Function
- Knowledge of Action
- Knowledge of Serial Order

Production System

Knowledge of Action in Sensorimotor Form:
- Attention at Key Points
- Action Programs

Mechanisms for Movement Control
- Environment
- Muscle Collectives

Figure 2: Roy and Square’s cognitive model of limb praxis, adapted from Roy and Square (1985).
Figure 3: A cognitive neuropsychological model of limb praxis. The three components on the right are interchangeable with the empty box in the complete model on the left. Under a) Rothi et al.’s original model of limb praxis. Under b) the previous model revised by Cubelli et al. and under c) the model extended by Buxbaum et al. For a detailed description see the text. Adapted respectively from Rothi et al. (1997), Cubelli et al. (2000) and Buxbaum et al. (2000).
Figure 4: An example of the abnormal kinematics of an apraxic movement. A patient with left ischemic lesions was tested in a study of imitation of meaningless gestures. The stimulus to imitate for this movement is shown under a) and represents a hand posture relative to the head. Under b), the movement times of the patient are longer than those of a matched normal subject (including replacement of the hand in the initial condition). Under c), the trajectory of the shoulder flexion-extension joint angle of the patient (shown in solid line) contains several irregularities which are the result from multiple hesitations and changes of directions, whereas the matched normal subject shoulder flexion-extension trajectory (dashed line) is smooth. The speed profile of the patient (solid line) is shown under d) and contains multiple peaks with reduced maximum velocities that contrast with the simple bell-shaped velocity profile of the matched normal subject (dashed line).
Figure 5: Schema of the monkey brain areas and their connectivity. Adapted from (Wise et al., 1997).